

(19) World Intellectual Property  
Organization  
International Bureau



102 (c)  
10 1459.834  
(43) International Publication Date  
12 August 2004 (12.08.2004)

PCT

(10) International Publication Number  
**WO 2004/066948 A2**

(51) International Patent Classification<sup>7</sup>:**A61K**

(74) Agents: SHAYESTEH, Laleh et al.; Exelixis, Inc., 170 Harbor Way, P.O. Box 511, South San Francisco, 5 94083-0511 (US).

(21) International Application Number:

PCT/US2004/002338

(81) Designated States (*unless otherwise indicated, for every kind of national protection available*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(30) Priority Data:

60/443,484	29 January 2003 (29.01.2003)	US
60/447,358	11 February 2003 (11.02.2003)	US
60/461,789	10 April 2003 (10.04.2003)	US
60/470,684	14 May 2003 (14.05.2003)	US
60/479,650	19 June 2003 (19.06.2003)	US

(71) Applicant (*for all designated States except US*): EX-ELIXIS INC. [US/US]; 170 Harbor Way, P.O. Box 511, South San Francisco, 5 94083-0511 (US).

(72) Inventors; and

(75) Inventors/Applicants (*for US only*): GENDREAU, Steven, Brian [US/US]; 2801 Turk Street, #103, San Francisco, CA 94118 (US). MORABLANCO, Eva, Lorena [US/US]; 145 Manor Drive, San Francisco, CA 94127 (US). LICKTEIG, Kim [US/US]; 1921 Grove Street, San Francisco, CA 94117 (US). ZHANG, HaiGuang [US/US]; 4833 El Grande Place, El Sobrante, CA 94083 (US).(84) Designated States (*unless otherwise indicated, for every kind of regional protection available*): ARIGO (BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

## Published:

— without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

**WO 2004/066948 A2**

(54) Title: MAPCAXS AS MODIFIERS OF THE APC AND AXIN PATHWAYS AND METHODS OF USE

(57) Abstract: Human MAPCAX genes are identified as modulators of the APC and axin pathways, and thus are therapeutic targets for disorders associated with defective APC and axin function. Methods for identifying modulators of APC and axin, comprising screening for agents that modulate the activity of MAPCAX are provided.

**MAPCAXs AS MODIFIERS OF THE APC AND AXIN PATHWAYS  
AND METHODS OF USE**

**REFERENCE TO RELATED APPLICATIONS**

5        This application claims priority to U.S. provisional patent applications 60/443,484 filed 1/29/2003, 60/447,358 filed 2/11/2003, 60/461,789 filed 4/10/2003, 60/470,684 filed 5/14/2003, and 60/479,650 filed 6/19/2003. The contents of the prior applications are hereby incorporated in their entirety.

10

**BACKGROUND OF THE INVENTION**

Deregulation of beta-catenin signaling is a frequent and early event in the development of a variety of human tumors, including colon cancer, melanoma, ovarian cancer, and prostate cancer. Activation of beta-catenin signaling can occur in tumor cells by loss-of-function mutations in the tumor suppressor genes APC (adenomatus polyposis coli protein) or Axin, as well as by gain-of-function mutations in the oncogene beta-catenin itself. APC, Axin, and beta-catenin normally bind to each other, as well as to the serine/threonine kinase GSK3-beta. Assembly of this degradation complex allows GSK3-beta to phosphorylate beta-catenin, which leads to beta-catenin ubiquitination and degradation by the proteasome. In the absence of APC or Axin activity, beta-catenin protein becomes stabilized and accumulates in the nucleus where it acts as a transcriptional co-activator with TCF for the induction of target genes, including the cell cycle regulators cyclin D1 and c-Myc.

The *C. elegans* gene *pry-1* is the structural and functional ortholog of vertebrate Axin (Korswagen HC et al. (2002) *Genes Dev.* 16:1291-302). PRY-1 is predicted to contain conserved RGS and DIX domains that, in Axin, bind APC and Dishevelled, respectively. Overexpression of the *C. elegans* *pry-1* gene in zebrafish can fully rescue the mutant phenotype of *masterblind*, the zebrafish Axin1 mutation. *pry-1* loss-of-function mutations produce several phenotypes that appear to result from increased beta-catenin signaling (Gleason JE et al. (2002) *Genes Dev.* 16:1281-90; Korswagen et al., *supra*). We find that the temperature-sensitive, reduction-of-function *pry-1* mutant *mu38* grown at 25°C produces a multivulva (Muv) phenotype in which approximately 30% of animals are induced to form ectopic vulvae. The *pry-1* Muv mutant phenotype is suppressed by RNAi-mediated inactivation the beta-catenin ortholog *bar-1* and the TCF ortholog *pop-1*. The Muv phenotype can also be generated by gain-of-function mutations

in *bar-1*/beta-catenin that eliminate the consensus GSK3-beta phosphorylation sites and are predicted to prevent Axin-mediated degradation of BAR-1.

The *C. elegans* gene product APR-1 shows significant structural similarity to human APC and can bind to both BAR-1/beta-catenin and PRY-1/Axin (Rocheleau et al. 5 (1997), Cell, Vol. 90, 707-716; Natarajan et al. (2001), Genetics, Vol. 159, 159-172; Korswagen et al., *supra*).

The ability to manipulate the genomes of model organisms such as *C. elegans* provides a powerful means to analyze biochemical processes that, due to significant evolutionary conservation, have direct relevance to more complex vertebrate organisms.

10 Due to a high level of gene and pathway conservation, the strong similarity of cellular processes, and the functional conservation of genes between these model organisms and mammals, identification of the involvement of novel genes in particular pathways and their functions in such model organisms can directly contribute to the understanding of the correlative pathways and methods of modulating them in mammals (see, for example,

15 Dulubova I, et al, J Neurochem 2001 Apr;77(1):229-38; Cai T, et al., Diabetologia 2001 Jan;44(1):81-8; Pasquinelli AE, et al., Nature. 2000 Nov 2;408(6808):37-8; Ivanov IP, et al., EMBO J 2000 Apr 17;19(8):1907-17; Vajo Z et al., Mamm Genome 1999 Oct;10(10):1000-4). For example, a genetic screen can be carried out in an invertebrate model organism having underexpression (e.g. knockout) or overexpression of a gene

20 (referred to as a "genetic entry point") that yields a visible phenotype. Additional genes are mutated in a random or targeted manner. When a gene mutation changes the original phenotype caused by the mutation in the genetic entry point, the gene is identified as a "modifier" involved in the same or overlapping pathway as the genetic entry point. When the genetic entry point is an ortholog of a human gene implicated in a disease pathway,

25 such as APC and axin, modifier genes can be identified that may be attractive candidate targets for novel therapeutics.

All references cited herein, including patents, patent applications, publications, and sequence information in referenced Genbank identifier numbers, are incorporated herein in their entireties.

30

## SUMMARY OF THE INVENTION

We have discovered genes that modify the APC and axin pathways in *C. elegans*, and identified their human orthologs, hereinafter referred to as modifier of APC and Axin (MAPCAX). The invention provides methods for utilizing these APC and axin modifier

genes and polypeptides to identify MAPCAX-modulating agents that are candidate therapeutic agents that can be used in the treatment of disorders associated with defective or impaired APC and axin function and/or MAPCAX function. Preferred MAPCAX-modulating agents specifically bind to MAPCAX polypeptides and restore APC and axin function. Other preferred MAPCAX-modulating agents are nucleic acid modulators such as antisense oligomers and RNAi that repress MAPCAX gene expression or product activity by, for example, binding to and inhibiting the respective nucleic acid (i.e. DNA or mRNA).

MAPCAX modulating agents may be evaluated by any convenient *in vitro* or *in vivo* assay for molecular interaction with a MAPCAX polypeptide or nucleic acid. In one embodiment, candidate MAPCAX modulating agents are tested with an assay system comprising a MAPCAX polypeptide or nucleic acid. Agents that produce a change in the activity of the assay system relative to controls are identified as candidate APC and axin modulating agents. The assay system may be cell-based or cell-free. MAPCAX-modulating agents include MAPCAX related proteins (e.g. dominant negative mutants, and biotherapeutics); MAPCAX -specific antibodies; MAPCAX -specific antisense oligomers and other nucleic acid modulators; and chemical agents that specifically bind to or interact with MAPCAX or compete with MAPCAX binding partner (e.g. by binding to a MAPCAX binding partner). In one specific embodiment, a small molecule modulator is identified using a binding assay. In specific embodiments, the screening assay system is selected from an apoptosis assay, a cell proliferation assay, an angiogenesis assay, and a hypoxic induction assay.

In another embodiment, candidate APC and axin pathways modulating agents are further tested using a second assay system that detects changes in the APC and axin pathways, such as angiogenic, apoptotic, or cell proliferation changes produced by the originally identified candidate agent or an agent derived from the original agent. The second assay system may use cultured cells or non-human animals. In specific embodiments, the secondary assay system uses non-human animals, including animals predetermined to have a disease or disorder implicating the APC and axin pathways, such as an angiogenic, apoptotic, or cell proliferation disorder (e.g. cancer).

The invention further provides methods for modulating the MAPCAX function and/or the APC and axin pathways in a mammalian cell by contacting the mammalian cell with an agent that specifically binds a MAPCAX polypeptide or nucleic acid. The agent may be a small molecule modulator, a nucleic acid modulator, or an antibody and may be

administered to a mammalian animal predetermined to have a pathology associated with the APC and axin pathways.

## DETAILED DESCRIPTION OF THE INVENTION

5       Genetic screens were designed to identify modifiers of the axin and APC pathway in *C. elegans*. The function of *apr-1* was depleted by RNAi in a *pry-1* hypomorphic allele *mu38*. Various specific genes were then silenced by RNA inhibition (RNAi). Methods for using RNAi to silence genes in *C. elegans* are known in the art (Fire A, et al., 1998 Nature 391:806-811; Fire, A. Trends Genet. 15, 358-363 (1999); WO9932619). Genes causing altered phenotypes in the worms were identified as modifiers of the APC and axin pathways. Modifiers of particular interest, were identified followed by identification of their orthologs. Accordingly, vertebrate orthologs of these modifiers, and preferably the human orthologs, MAPCAX genes (i.e., nucleic acids and polypeptides) are attractive drug targets for the treatment of pathologies associated with a defective APC and axin 10 signaling pathway, such as cancer. Table 1 (Example II) lists the modifiers and their 15 orthologs.

In vitro and in vivo methods of assessing MAPCAX function are provided herein. Modulation of the MAPCAX or their respective binding partners is useful for understanding the association of the APC and axin pathways and their members in normal 20 and disease conditions and for developing diagnostics and therapeutic modalities for APC and axin related pathologies. MAPCAX-modulating agents that act by inhibiting or enhancing MAPCAX expression, directly or indirectly, for example, by affecting a MAPCAX function such as enzymatic (e.g., catalytic) or binding activity, can be identified using methods provided herein. MAPCAX modulating agents are useful in 25 diagnosis, therapy and pharmaceutical development.

### Nucleic acids and polypeptides of the invention

Sequences related to MAPCAX nucleic acids and polypeptides that can be used in the invention are disclosed in Genbank (referenced by Genbank identifier (GI) or RefSeq 30 number), shown in Table 1 and in the sequence listing.

The term “MAPCAX polypeptide” refers to a full-length MAPCAX protein or a functionally active fragment or derivative thereof. A “functionally active” MAPCAX fragment or derivative exhibits one or more functional activities associated with a full-length, wild-type MAPCAX protein, such as antigenic or immunogenic activity,

enzymatic activity, ability to bind natural cellular substrates, etc. The functional activity of MAPCAX proteins, derivatives and fragments can be assayed by various methods known to one skilled in the art (Current Protocols in Protein Science (1998) Coligan *et al.*, eds., John Wiley & Sons, Inc., Somerset, New Jersey) and as further discussed below. In 5 one embodiment, a functionally active MAPCAX polypeptide is a MAPCAX derivative capable of rescuing defective endogenous MAPCAX activity, such as in cell based or animal assays; the rescuing derivative may be from the same or a different species. For purposes herein, functionally active fragments also include those fragments that comprise 10 one or more structural domains of a MAPCAX, such as a binding domain. Protein domains can be identified using the PFAM program (Bateman A., et al., Nucleic Acids Res, 1999, 27:260-2). Methods for obtaining MAPCAX polypeptides are also further described below. In some embodiments, preferred fragments are functionally active, 15 domain-containing fragments comprising at least 25 contiguous amino acids, preferably at least 50, more preferably 75, and most preferably at least 100 contiguous amino acids of a MAPCAX. In further preferred embodiments, the fragment comprises the entire functionally active domain.

The term "MAPCAX nucleic acid" refers to a DNA or RNA molecule that encodes 20 a MAPCAX polypeptide. Preferably, the MAPCAX polypeptide or nucleic acid or fragment thereof is from a human, but can also be an ortholog, or derivative thereof with at least 70% sequence identity, preferably at least 80%, more preferably 85%, still more preferably 90%, and most preferably at least 95% sequence identity with human MAPCAX. Methods of identifying orthologs are known in the art. Normally, orthologs in 25 different species retain the same function, due to presence of one or more protein motifs and/or 3-dimensional structures. Orthologs are generally identified by sequence homology analysis, such as BLAST analysis, usually using protein bait sequences. Sequences are assigned as a potential ortholog if the best hit sequence from the forward BLAST result retrieves the original query sequence in the reverse BLAST (Huynen MA and Bork P, Proc Natl Acad Sci (1998) 95:5849-5856; Huynen MA *et al.*, Genome Research (2000) 10:1204-1210). Programs for multiple sequence alignment, such as CLUSTAL 30 (Thompson JD *et al*, 1994, Nucleic Acids Res 22:4673-4680) may be used to highlight conserved regions and/or residues of orthologous proteins and to generate phylogenetic trees. In a phylogenetic tree representing multiple homologous sequences from diverse species (e.g., retrieved through BLAST analysis), orthologous sequences from two species generally appear closest on the tree with respect to all other sequences from these two

species. Structural threading or other analysis of protein folding (e.g., using software by ProCeryon, Biosciences, Salzburg, Austria) may also identify potential orthologs. In evolution, when a gene duplication event follows speciation, a single gene in one species, such as *C. elegans*, may correspond to multiple genes (paralogs) in another, such as

5 human. As used herein, the term "orthologs" encompasses paralogs. As used herein, "percent (%) sequence identity" with respect to a subject sequence, or a specified portion of a subject sequence, is defined as the percentage of nucleotides or amino acids in the candidate derivative sequence identical with the nucleotides or amino acids in the subject sequence (or specified portion thereof), after aligning the sequences and introducing gaps,

10 if necessary to achieve the maximum percent sequence identity, as generated by the program WU-BLAST-2.0a19 (Altschul *et al.*, J. Mol. Biol. (1997) 215:403-410) with all the search parameters set to default values. The HSP S and HSP S2 parameters are dynamic values and are established by the program itself depending upon the composition of the particular sequence and composition of the particular database against which the

15 sequence of interest is being searched. A % identity value is determined by the number of matching identical nucleotides or amino acids divided by the sequence length for which the percent identity is being reported. "Percent (%) amino acid sequence similarity" is determined by doing the same calculation as for determining % amino acid sequence identity, but including conservative amino acid substitutions in addition to identical amino

20 acids in the computation.

A conservative amino acid substitution is one in which an amino acid is substituted for another amino acid having similar properties such that the folding or activity of the protein is not significantly affected. Aromatic amino acids that can be substituted for each other are phenylalanine, tryptophan, and tyrosine; interchangeable hydrophobic amino acids are leucine, isoleucine, methionine, and valine; interchangeable polar amino acids are glutamine and asparagine; interchangeable basic amino acids are arginine, lysine and histidine; interchangeable acidic amino acids are aspartic acid and glutamic acid; and interchangeable small amino acids are alanine, serine, threonine, cysteine and glycine.

Alternatively, an alignment for nucleic acid sequences is provided by the local homology algorithm of Smith and Waterman (Smith and Waterman, 1981, Advances in Applied Mathematics 2:482-489; database: European Bioinformatics Institute; Smith and Waterman, 1981, J. of Molec.Biol., 147:195-197; Nicholas *et al.*, 1998, "A Tutorial on Searching Sequence Databases and Sequence Scoring Methods" ([www.psc.edu](http://www.psc.edu)) and references cited therein.; W.R. Pearson, 1991, Genomics 11:635-650). This algorithm can

be applied to amino acid sequences by using the scoring matrix developed by Dayhoff (Dayhoff: Atlas of Protein Sequences and Structure, M. O. Dayhoff ed., 5 suppl. 3:353-358, National Biomedical Research Foundation, Washington, D.C., USA), and normalized by Gribskov (Gribskov 1986 Nucl. Acids Res. 14(6):6745-6763). The Smith-Waterman 5 algorithm may be employed where default parameters are used for scoring (for example, gap open penalty of 12, gap extension penalty of two). From the data generated, the "Match" value reflects "sequence identity."

Derivative nucleic acid molecules of the subject nucleic acid molecules include sequences that hybridize to the nucleic acid sequence of a MAPCAX. The stringency of 10 hybridization can be controlled by temperature, ionic strength, pH, and the presence of denaturing agents such as formamide during hybridization and washing. Conditions routinely used are set out in readily available procedure texts (e.g., Current Protocol in Molecular Biology, Vol. 1, Chap. 2.10, John Wiley & Sons, Publishers (1994); Sambrook 15 et al., Molecular Cloning, Cold Spring Harbor (1989)). In some embodiments, a nucleic acid molecule of the invention is capable of hybridizing to a nucleic acid molecule containing the nucleotide sequence of a MAPCAX under high stringency hybridization conditions that are: prehybridization of filters containing nucleic acid for 8 hours to overnight at 65° C in a solution comprising 6X single strength citrate (SSC) (1X SSC is 0.15 M NaCl, 0.015 M Na citrate; pH 7.0), 5X Denhardt's solution, 0.05% sodium 20 pyrophosphate and 100 µg/ml herring sperm DNA; hybridization for 18-20 hours at 65° C in a solution containing 6X SSC, 1X Denhardt's solution, 100 µg/ml yeast tRNA and 0.05% sodium pyrophosphate; and washing of filters at 65° C for 1h in a solution containing 0.1X SSC and 0.1% SDS (sodium dodecyl sulfate).

In other embodiments, moderately stringent hybridization conditions are used that 25 are: pretreatment of filters containing nucleic acid for 6 h at 40° C in a solution containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH7.5), 5mM EDTA, 0.1% PVP, 0.1% Ficoll, 1% BSA, and 500 µg/ml denatured salmon sperm DNA; hybridization for 18-20h at 40° C in a solution containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH7.5), 5mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.2% BSA, 100 µg/ml salmon sperm DNA, and 30 10% (wt/vol) dextran sulfate; followed by washing twice for 1 hour at 55° C in a solution containing 2X SSC and 0.1% SDS.

Alternatively, low stringency conditions can be used that are: incubation for 8 hours to overnight at 37° C in a solution comprising 20% formamide, 5 x SSC, 50 mM sodium phosphate (pH 7.6), 5X Denhardt's solution, 10% dextran sulfate, and 20 µg/ml

denatured sheared salmon sperm DNA; hybridization in the same buffer for 18 to 20 hours; and washing of filters in 1 x SSC at about 37° C for 1 hour.

### Isolation, Production, Expression, and Mis-expression of MAPCAX Nucleic Acids and Polypeptides

MAPCAX nucleic acids and polypeptides are useful for identifying and testing agents that modulate MAPCAX function and for other applications related to the involvement of MAPCAX in the APC and axin pathways. MAPCAX nucleic acids and derivatives and orthologs thereof may be obtained using any available method. For instance, techniques for isolating cDNA or genomic DNA sequences of interest by screening DNA libraries or by using polymerase chain reaction (PCR) are well known in the art. In general, the particular use for the protein will dictate the particulars of expression, production, and purification methods. For instance, production of proteins for use in screening for modulating agents may require methods that preserve specific biological activities of these proteins, whereas production of proteins for antibody generation may require structural integrity of particular epitopes. Expression of proteins to be purified for screening or antibody production may require the addition of specific tags (*e.g.*, generation of fusion proteins). Overexpression of a MAPCAX protein for assays used to assess MAPCAX function, such as involvement in cell cycle regulation or hypoxic response, may require expression in eukaryotic cell lines capable of these cellular activities. Techniques for the expression, production, and purification of proteins are well known in the art; any suitable means therefore may be used (*e.g.*, Higgins SJ and Hames BD (eds.) *Protein Expression: A Practical Approach*, Oxford University Press Inc., New York 1999; Stanbury PF et al., *Principles of Fermentation Technology*, 2<sup>nd</sup> edition, Elsevier Science, New York, 1995; Doonan S (ed.) *Protein Purification Protocols*, Humana Press, New Jersey, 1996; Coligan JE et al, *Current Protocols in Protein Science* (eds.), 1999, John Wiley & Sons, New York). In particular embodiments, recombinant MAPCAX is expressed in a cell line known to have defective APC or axin function. The recombinant cells are used in cell-based screening assay systems of the invention, as described further below.

The nucleotide sequence encoding a MAPCAX polypeptide can be inserted into any appropriate expression vector. The necessary transcriptional and translational signals, including promoter/enhancer element, can derive from the native MAPCAX gene and/or its flanking regions or can be heterologous. A variety of host-vector expression systems

may be utilized, such as mammalian cell systems infected with virus (e.g. vaccinia virus, adenovirus, etc.); insect cell systems infected with virus (e.g. baculovirus); microorganisms such as yeast containing yeast vectors, or bacteria transformed with bacteriophage, plasmid, or cosmid DNA. An isolated host cell strain that modulates the expression of, modifies, and/or specifically processes the gene product may be used.

To detect expression of the MAPCAX gene product, the expression vector can comprise a promoter operably linked to a MAPCAX gene nucleic acid, one or more origins of replication, and, one or more selectable markers (e.g. thymidine kinase activity, resistance to antibiotics, etc.). Alternatively, recombinant expression vectors can be identified by assaying for the expression of the MAPCAX gene product based on the physical or functional properties of the MAPCAX protein in *in vitro* assay systems (e.g. immunoassays).

The MAPCAX protein, fragment, or derivative may be optionally expressed as a fusion, or chimeric protein product (i.e. it is joined via a peptide bond to a heterologous protein sequence of a different protein), for example to facilitate purification or detection. A chimeric product can be made by ligating the appropriate nucleic acid sequences encoding the desired amino acid sequences to each other using standard methods and expressing the chimeric product. A chimeric product may also be made by protein synthetic techniques, e.g. by use of a peptide synthesizer (Hunkapiller *et al.*, Nature (1984) 20 310:105-111).

Once a recombinant cell that expresses the MAPCAX gene sequence is identified, the gene product can be isolated and purified using standard methods (e.g. ion exchange, affinity, and gel exclusion chromatography; centrifugation; differential solubility; electrophoresis). Alternatively, native MAPCAX proteins can be purified from natural sources, by standard methods (e.g. immunoaffinity purification). Once a protein is obtained, it may be quantified and its activity measured by appropriate methods, such as immunoassay, bioassay, or other measurements of physical properties, such as crystallography.

The methods of this invention may also use cells that have been engineered for altered expression (mis-expression) of MAPCAX or other genes associated with the APC and axin pathways. As used herein, mis-expression encompasses ectopic expression, over-expression, under-expression, and non-expression (e.g. by gene knock-out or blocking expression that would otherwise normally occur).

### Genetically modified animals

Animal models that have been genetically modified to alter MAPCAX expression may be used in *in vivo* assays to test for activity of a candidate APC and axin modulating agent, or to further assess the role of MAPCAX in a APC and axin pathways process such as apoptosis or cell proliferation. Preferably, the altered MAPCAX expression results in a detectable phenotype, such as decreased or increased levels of cell proliferation, angiogenesis, or apoptosis compared to control animals having normal MAPCAX expression. The genetically modified animal may additionally have altered APC and axin expression (e.g. APC and axin knockout). Preferred genetically modified animals are mammals such as primates, rodents (preferably mice or rats), among others. Preferred non-mammalian species include zebrafish, *C. elegans*, and *Drosophila*. Preferred genetically modified animals are transgenic animals having a heterologous nucleic acid sequence present as an extrachromosomal element in a portion of its cells, i.e. mosaic animals (see, for example, techniques described by Jakobovits, 1994, Curr. Biol. 4:761-763.) or stably integrated into its germ line DNA (i.e., in the genomic sequence of most or all of its cells). Heterologous nucleic acid is introduced into the germ line of such transgenic animals by genetic manipulation of, for example, embryos or embryonic stem cells of the host animal.

Methods of making transgenic animals are well-known in the art (for transgenic mice see Brinster et al., Proc. Nat. Acad. Sci. USA 82: 4438-4442 (1985), U.S. Pat. Nos. 4,736,866 and 4,870,009, both by Leder et al., U.S. Pat. No. 4,873,191 by Wagner et al., and Hogan, B., Manipulating the Mouse Embryo, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., (1986); for particle bombardment see U.S. Pat. No., 4,945,050, by Sandford et al.; for transgenic *Drosophila* see Rubin and Spradling, Science (1982) 218:348-53 and U.S. Pat. No. 4,670,388; for transgenic insects see Berghammer A.J. et al., A Universal Marker for Transgenic Insects (1999) Nature 402:370-371; for transgenic Zebrafish see Lin S., Transgenic Zebrafish, Methods Mol Biol. (2000);136:375-3830); for microinjection procedures for fish, amphibian eggs and birds see Houdebine and Chourrout, Experientia (1991) 47:897-905; for transgenic rats see Hammer et al., Cell (1990) 63:1099-1112; and for culturing of embryonic stem (ES) cells and the subsequent production of transgenic animals by the introduction of DNA into ES cells using methods such as electroporation, calcium phosphate/DNA precipitation and direct injection see, e.g., Teratocarcinomas and Embryonic Stem Cells, A Practical Approach, E. J. Robertson, ed., IRL Press (1987)). Clones of the nonhuman transgenic animals can be produced

according to available methods (see Wilmut, I. *et al.* (1997) Nature 385:810-813; and PCT International Publication Nos. WO 97/07668 and WO 97/07669).

In one embodiment, the transgenic animal is a "knock-out" animal having a heterozygous or homozygous alteration in the sequence of an endogenous MAPCAX gene 5 that results in a decrease of MAPCAX function, preferably such that MAPCAX expression is undetectable or insignificant. Knock-out animals are typically generated by homologous recombination with a vector comprising a transgene having at least a portion of the gene to be knocked out. Typically a deletion, addition or substitution has been introduced into the transgene to functionally disrupt it. The transgene can be a human 10 gene (e.g., from a human genomic clone) but more preferably is an ortholog of the human gene derived from the transgenic host species. For example, a mouse MAPCAX gene is used to construct a homologous recombination vector suitable for altering an endogenous MAPCAX gene in the mouse genome. Detailed methodologies for homologous recombination in mice are available (see Capecchi, Science (1989) 244:1288-1292; Joyner 15 *et al.*, Nature (1989) 338:153-156). Procedures for the production of non-rodent transgenic mammals and other animals are also available (Houdebine and Chourrout, *supra*; Pursel *et al.*, Science (1989) 244:1281-1288; Simms *et al.*, Bio/Technology (1988) 6:179-183). In a preferred embodiment, knock-out animals, such as mice harboring a knockout of a specific gene, may be used to produce antibodies against the human 20 counterpart of the gene that has been knocked out (Claesson MH *et al.*, (1994) Scan J Immunol 40:257-264; Declerck PJ *et al.*, (1995) J Biol Chem. 270:8397-400).

In another embodiment, the transgenic animal is a "knock-in" animal having an alteration in its genome that results in altered expression (e.g., increased (including ectopic) or decreased expression) of the MAPCAX gene, e.g., by introduction of 25 additional copies of MAPCAX, or by operatively inserting a regulatory sequence that provides for altered expression of an endogenous copy of the MAPCAX gene. Such regulatory sequences include inducible, tissue-specific, and constitutive promoters and enhancer elements. The knock-in can be homozygous or heterozygous.

Transgenic nonhuman animals can also be produced that contain selected systems 30 allowing for regulated expression of the transgene. One example of such a system that may be produced is the cre/loxP recombinase system of bacteriophage P1 (Lakso *et al.*, PNAS (1992) 89:6232-6236; U.S. Pat. No. 4,959,317). If a cre/loxP recombinase system is used to regulate expression of the transgene, animals containing transgenes encoding both the Cre recombinase and a selected protein are required. Such animals can be

provided through the construction of "double" transgenic animals, e.g., by mating two transgenic animals, one containing a transgene encoding a selected protein and the other containing a transgene encoding a recombinase. Another example of a recombinase system is the FLP recombinase system of *Saccharomyces cerevisiae* (O'Gorman et al. 5 (1991) *Science* 251:1351-1355; U.S. Pat. No. 5,654,182). In a preferred embodiment, both Cre-LoxP and Flp-Frt are used in the same system to regulate expression of the transgene, and for sequential deletion of vector sequences in the same cell (Sun X et al (2000) *Nat Genet* 25:83-6).

The genetically modified animals can be used in genetic studies to further elucidate 10 the APC and axin pathways, as animal models of disease and disorders implicating defective APC and axin function, and for *in vivo* testing of candidate therapeutic agents, such as those identified in screens described below. The candidate therapeutic agents are administered to a genetically modified animal having altered MAPCAX function and phenotypic changes are compared with appropriate control animals such as genetically 15 modified animals that receive placebo treatment, and/or animals with unaltered MAPCAX expression that receive candidate therapeutic agent.

In addition to the above-described genetically modified animals having altered MAPCAX function, animal models having defective APC and axin function (and otherwise normal MAPCAX function), can be used in the methods of the present 20 invention. For example, a APC and axin knockout mouse can be used to assess, *in vivo*, the activity of a candidate APC and axin modulating agent identified in one of the *in vitro* assays described below. Preferably, the candidate APC and axin modulating agent when administered to a model system with cells defective in APC and axin function, produces a detectable phenotypic change in the model system indicating that the APC and axin 25 function is restored, i.e., the cells exhibit normal cell cycle progression.

#### Modulating Agents

The invention provides methods to identify agents that interact with and/or modulate the function of MAPCAX and/or the APC and axin pathways. Modulating 30 agents identified by the methods are also part of the invention. Such agents are useful in a variety of diagnostic and therapeutic applications associated with the APC and axin pathways, as well as in further analysis of the MAPCAX protein and its contribution to the APC and axin pathways. Accordingly, the invention also provides methods for

modulating the APC and axin pathways comprising the step of specifically modulating MAPCAX activity by administering a MAPCAX-interacting or -modulating agent.

As used herein, an "MAPCAX-modulating agent" is any agent that modulates MAPCAX function, for example, an agent that interacts with MAPCAX to inhibit or enhance MAPCAX activity or otherwise affect normal MAPCAX function. MAPCAX function can be affected at any level, including transcription, protein expression, protein localization, and cellular or extra-cellular activity. In a preferred embodiment, the MAPCAX - modulating agent specifically modulates the function of the MAPCAX. The phrases "specific modulating agent", "specifically modulates", etc., are used herein to refer to modulating agents that directly bind to the MAPCAX polypeptide or nucleic acid, and preferably inhibit, enhance, or otherwise alter, the function of the MAPCAX. These phrases also encompass modulating agents that alter the interaction of the MAPCAX with a binding partner, substrate, or cofactor (e.g. by binding to a binding partner of a MAPCAX, or to a protein/binding partner complex, and altering MAPCAX function). In a further preferred embodiment, the MAPCAX- modulating agent is a modulator of the APC and axin pathways (e.g. it restores and/or upregulates APC and axin function) and thus is also an APC and axin-modulating agent.

Preferred MAPCAX-modulating agents include small molecule compounds; MAPCAX-interacting proteins, including antibodies and other biotherapeutics; and nucleic acid modulators such as antisense and RNA inhibitors. The modulating agents may be formulated in pharmaceutical compositions, for example, as compositions that may comprise other active ingredients, as in combination therapy, and/or suitable carriers or excipients. Techniques for formulation and administration of the compounds may be found in "Remington's Pharmaceutical Sciences" Mack Publishing Co., Easton, PA, 19<sup>th</sup> edition.

#### **Small molecule modulators**

Small molecules are often preferred to modulate function of proteins with enzymatic function, and/or containing protein interaction domains. Chemical agents, referred to in the art as "small molecule" compounds are typically organic, non-peptide molecules, having a molecular weight up to 10,000, preferably up to 5,000, more preferably up to 1,000, and most preferably up to 500 daltons. This class of modulators includes chemically synthesized molecules, for instance, compounds from combinatorial chemical libraries. Synthetic compounds may be rationally designed or identified based

on known or inferred properties of the MAPCAX protein or may be identified by screening compound libraries. Alternative appropriate modulators of this class are natural products, particularly secondary metabolites from organisms such as plants or fungi, which can also be identified by screening compound libraries for MAPCAX-modulating activity. Methods for generating and obtaining compounds are well known in the art (Schreiber SL, Science (2000) 151: 1964-1969; Radmann J and Gunther J, Science (2000) 151:1947-1948).

Small molecule modulators identified from screening assays, as described below, can be used as lead compounds from which candidate clinical compounds may be designed, optimized, and synthesized. Such clinical compounds may have utility in treating pathologies associated with the APC and axin pathways. The activity of candidate small molecule modulating agents may be improved several-fold through iterative secondary functional validation, as further described below, structure determination, and candidate modulator modification and testing. Additionally, candidate clinical compounds are generated with specific regard to clinical and pharmacological properties. For example, the reagents may be derivatized and re-screened using *in vitro* and *in vivo* assays to optimize activity and minimize toxicity for pharmaceutical development.

### Protein Modulators

Specific MAPCAX-interacting proteins are useful in a variety of diagnostic and therapeutic applications related to the APC and axin pathways and related disorders, as well as in validation assays for other MAPCAX-modulating agents. In a preferred embodiment, MAPCAX-interacting proteins affect normal MAPCAX function, including transcription, protein expression, protein localization, and cellular or extra-cellular activity. In another embodiment, MAPCAX-interacting proteins are useful in detecting and providing information about the function of MAPCAX proteins, as is relevant to APC and axin related disorders, such as cancer (e.g., for diagnostic means).

A MAPCAX-interacting protein may be endogenous, i.e. one that naturally interacts genetically or biochemically with a MAPCAX, such as a member of the MAPCAX pathway that modulates MAPCAX expression, localization, and/or activity. MAPCAX-modulators include dominant negative forms of MAPCAX-interacting proteins and of MAPCAX proteins themselves. Yeast two-hybrid and variant screens offer preferred methods for identifying endogenous MAPCAX-interacting proteins (Finley, R. L. et al. (1996) in DNA Cloning-Expression Systems: A Practical Approach, eds. Glover

D. & Hames B. D (Oxford University Press, Oxford, England), pp. 169-203; Fashema SF et al., Gene (2000) 250:1-14; Drees BL Curr Opin Chem Biol (1999) 3:64-70; Vidal M and Legrain P Nucleic Acids Res (1999) 27:919-29; and U.S. Pat. No. 5,928,868). Mass spectrometry is an alternative preferred method for the elucidation of protein complexes  
5 (reviewed in, e.g., Pandley A and Mann M, Nature (2000) 405:837-846; Yates JR 3<sup>rd</sup>, Trends Genet (2000) 16:5-8).

A MAPCAX-interacting protein may be an exogenous protein, such as a MAPCAX-specific antibody or a T-cell antigen receptor (see, e.g., Harlow and Lane (1988) Antibodies, A Laboratory Manual, Cold Spring Harbor Laboratory; Harlow and Lane (1999) Using antibodies: a laboratory manual. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press). MAPCAX antibodies are further discussed below.  
10

In preferred embodiments, a MAPCAX-interacting protein specifically binds a MAPCAX protein. In alternative preferred embodiments, a MAPCAX-modulating agent binds a MAPCAX substrate, binding partner, or cofactor.

15

### ***Antibodies***

In another embodiment, the protein modulator is a MAPCAX specific antibody agonist or antagonist. The antibodies have therapeutic and diagnostic utilities, and can be used in screening assays to identify MAPCAX modulators. The antibodies can also be  
20 used in dissecting the portions of the MAPCAX pathway responsible for various cellular responses and in the general processing and maturation of the MAPCAX.

Antibodies that specifically bind MAPCAX polypeptides can be generated using known methods. Preferably the antibody is specific to a mammalian ortholog of MAPCAX polypeptide, and more preferably, to human MAPCAX. Antibodies may be  
25 polyclonal, monoclonal (mAbs), humanized or chimeric antibodies, single chain antibodies, Fab fragments, F(ab').sub.2 fragments, fragments produced by a FAb expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above. Epitopes of MAPCAX which are particularly antigenic can be selected, for example, by routine screening of MAPCAX polypeptides for antigenicity or by  
30 applying a theoretical method for selecting antigenic regions of a protein (Hopp and Wood (1981), Proc. Nati. Acad. Sci. U.S.A. 78:3824-28; Hopp and Wood, (1983) Mol. Immunol. 20:483-89; Sutcliffe et al., (1983) Science 219:660-66) to the amino acid sequence of a MAPCAX. Monoclonal antibodies with affinities of  $10^8 \text{ M}^{-1}$  preferably  $10^9 \text{ M}^{-1}$  to  $10^{10} \text{ M}^{-1}$ , or stronger can be made by standard procedures as described (Harlow and Lane, *supra*;

Goding (1986) *Monoclonal Antibodies: Principles and Practice* (2d ed) Academic Press, New York; and U.S. Pat. Nos. 4,381,292; 4,451,570; and 4,618,577). Antibodies may be generated against crude cell extracts of MAPCAX or substantially purified fragments thereof. If MAPCAX fragments are used, they preferably comprise at least 10, and more 5 preferably, at least 20 contiguous amino acids of a MAPCAX protein. In a particular embodiment, MAPCAX-specific antigens and/or immunogens are coupled to carrier proteins that stimulate the immune response. For example, the subject polypeptides are covalently coupled to the keyhole limpet hemocyanin (KLH) carrier, and the conjugate is emulsified in Freund's complete adjuvant, which enhances the immune response. An 10 appropriate immune system such as a laboratory rabbit or mouse is immunized according to conventional protocols.

The presence of MAPCAX-specific antibodies is assayed by an appropriate assay such as a solid phase enzyme-linked immunosorbant assay (ELISA) using immobilized corresponding MAPCAX polypeptides. Other assays, such as radioimmunoassays or 15 fluorescent assays might also be used.

Chimeric antibodies specific to MAPCAX polypeptides can be made that contain different portions from different animal species. For instance, a human immunoglobulin constant region may be linked to a variable region of a murine mAb, such that the antibody derives its biological activity from the human antibody, and its binding 20 specificity from the murine fragment. Chimeric antibodies are produced by splicing together genes that encode the appropriate regions from each species (Morrison et al., Proc. Natl. Acad. Sci. (1984) 81:6851-6855; Neuberger et al., Nature (1984) 312:604-608; Takeda et al., Nature (1985) 31:452-454). Humanized antibodies, which are a form of chimeric antibodies, can be generated by grafting complementary-determining regions 25 (CDRs) (Carlos, T. M., J. M. Harlan. 1994. Blood 84:2068-2101) of mouse antibodies into a background of human framework regions and constant regions by recombinant DNA technology (Riechmann LM, et al., 1988 Nature 323: 323-327). Humanized antibodies contain ~10% murine sequences and ~90% human sequences, and thus further reduce or eliminate immunogenicity, while retaining the antibody specificities (Co MS, 30 and Queen C. 1991 Nature 351: 501-501; Morrison SL. 1992 Ann. Rev. Immun. 10:239-265). Humanized antibodies and methods of their production are well-known in the art (U.S. Pat. Nos. 5,530,101, 5,585,089, 5,693,762, and 6,180,370).

MAPCAX-specific single chain antibodies which are recombinant, single chain polypeptides formed by linking the heavy and light chain fragments of the Fv regions via

an amino acid bridge, can be produced by methods known in the art (U.S. Pat. No. 4,946,778; Bird, Science (1988) 242:423-426; Huston et al., Proc. Natl. Acad. Sci. USA (1988) 85:5879-5883; and Ward et al., Nature (1989) 334:544-546).

Other suitable techniques for antibody production involve in vitro exposure of 5 lymphocytes to the antigenic polypeptides or alternatively to selection of libraries of antibodies in phage or similar vectors (Huse et al., Science (1989) 246:1275-1281). As used herein, T-cell antigen receptors are included within the scope of antibody modulators (Harlow and Lane, 1988, *supra*).

The polypeptides and antibodies of the present invention may be used with or 10 without modification. Frequently, antibodies will be labeled by joining, either covalently or non-covalently, a substance that provides for a detectable signal, or that is toxic to cells that express the targeted protein (Menard S, et al., Int J. Biol Markers (1989) 4:131-134). A wide variety of labels and conjugation techniques are known and are reported extensively in both the scientific and patent literature. Suitable labels include 15 radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent moieties, fluorescent emitting lanthanide metals, chemiluminescent moieties, bioluminescent moieties, magnetic particles, and the like (U.S. Pat. Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241). Also, recombinant immunoglobulins may be produced (U.S. Pat. No. 4,816,567). Antibodies to cytoplasmic polypeptides may 20 be delivered and reach their targets by conjugation with membrane-penetrating toxin proteins (U.S. Pat. No. 6,086,900).

When used therapeutically in a patient, the antibodies of the subject invention are typically administered parenterally, when possible at the target site, or intravenously. The therapeutically effective dose and dosage regimen is determined by clinical studies. 25 Typically, the amount of antibody administered is in the range of about 0.1 mg/kg –to about 10 mg/kg of patient weight. For parenteral administration, the antibodies are formulated in a unit dosage injectable form (e.g., solution, suspension, emulsion) in association with a pharmaceutically acceptable vehicle. Such vehicles are inherently nontoxic and non-therapeutic. Examples are water, saline, Ringer's solution, dextrose 30 solution, and 5% human serum albumin. Nonaqueous vehicles such as fixed oils, ethyl oleate, or liposome carriers may also be used. The vehicle may contain minor amounts of additives, such as buffers and preservatives, which enhance isotonicity and chemical stability or otherwise enhance therapeutic potential. The antibodies' concentrations in such vehicles are typically in the range of about 1 mg/ml to about 10 mg/ml.

Immunotherapeutic methods are further described in the literature (US Pat. No. 5,859,206; WO0073469).

### Nucleic Acid Modulators

- 5 Other preferred MAPCAX-modulating agents comprise nucleic acid molecules, such as antisense oligomers or double stranded RNA (dsRNA), which generally inhibit MAPCAX activity. Preferred nucleic acid modulators interfere with the function of the MAPCAX nucleic acid such as DNA replication, transcription, translocation of the MAPCAX RNA to the site of protein translation, translation of protein from the  
10 MAPCAX RNA, splicing of the MAPCAX RNA to yield one or more mRNA species, or catalytic activity which may be engaged in or facilitated by the MAPCAX RNA.

In one embodiment, the antisense oligomer is an oligonucleotide that is sufficiently complementary to a MAPCAX mRNA to bind to and prevent translation, preferably by binding to the 5' untranslated region. MAPCAX-specific antisense oligonucleotides, 15 preferably range from at least 6 to about 200 nucleotides. In some embodiments the oligonucleotide is preferably at least 10, 15, or 20 nucleotides in length. In other embodiments, the oligonucleotide is preferably less than 50, 40, or 30 nucleotides in length. The oligonucleotide can be DNA or RNA or a chimeric mixture or derivatives or modified versions thereof, single-stranded or double-stranded. The oligonucleotide can be 20 modified at the base moiety, sugar moiety, or phosphate backbone. The oligonucleotide may include other appending groups such as peptides, agents that facilitate transport across the cell membrane, hybridization-triggered cleavage agents, and intercalating agents.

In another embodiment, the antisense oligomer is a phosphothioate morpholino 25 oligomer (PMO). PMOs are assembled from four different morpholino subunits, each of which contain one of four genetic bases (A, C, G, or T) linked to a six-membered morpholine ring. Polymers of these subunits are joined by non-ionic phosphodiamidate intersubunit linkages. Details of how to make and use PMOs and other antisense oligomers are well known in the art (e.g. see WO99/18193; Probst JC, Antisense 30 Oligodeoxynucleotide and Ribozyme Design, Methods. (2000) 22(3):271-281; Summerton J, and Weller D. 1997 Antisense Nucleic Acid Drug Dev. :7:187-95; US Pat. No. 5,235,033; and US Pat No. 5,378,841).

Alternative preferred MAPCAX nucleic acid modulators are double-stranded RNA species mediating RNA interference (RNAi). RNAi is the process of sequence-specific,

post-transcriptional gene silencing in animals and plants, initiated by double-stranded RNA (dsRNA) that is homologous in sequence to the silenced gene. Methods relating to the use of RNAi to silence genes in *C. elegans*, *Drosophila*, plants, and humans are known in the art (Fire A, et al., 1998 *Nature* 391:806-811; Fire, A. *Trends Genet.* 15, 358-363 (1999); Sharp, P. A. *RNA interference* 2001. *Genes Dev.* 15, 485-490 (2001); Hammond, S. M., et al., *Nature Rev. Genet.* 2, 110-1119 (2001); Tuschl, T. *Chem. Biochem.* 2, 239-245 (2001); Hamilton, A. et al., *Science* 286, 950-952 (1999); Hammond, S. M., et al., *Nature* 404, 293-296 (2000); Zamore, P. D., et al., *Cell* 101, 25-33 (2000); Bernstein, E., et al., *Nature* 409, 363-366 (2001); Elbashir, S. M., et al., *Genes Dev.* 15, 188-200 (2001); WO0129058; WO9932619; Elbashir SM, et al., 2001 *Nature* 411:494-498).

Nucleic acid modulators are commonly used as research reagents, diagnostics, and therapeutics. For example, antisense oligonucleotides, which are able to inhibit gene expression with exquisite specificity, are often used to elucidate the function of particular genes (see, for example, U.S. Pat. No. 6,165,790). Nucleic acid modulators are also used, for example, to distinguish between functions of various members of a biological pathway. For example, antisense oligomers have been employed as therapeutic moieties in the treatment of disease states in animals and man and have been demonstrated in numerous clinical trials to be safe and effective (Milligan JF, et al, *Current Concepts in Antisense Drug Design*, *J Med Chem.* (1993) 36:1923-1937; Tonkinson JL et al., *Antisense Oligodeoxynucleotides as Clinical Therapeutic Agents*, *Cancer Invest.* (1996) 14:54-65). Accordingly, in one aspect of the invention, a MAPCAX-specific nucleic acid modulator is used in an assay to further elucidate the role of the MAPCAX in the APC and axin pathways, and/or its relationship to other members of the pathway. In another aspect of the invention, a MAPCAX-specific antisense oligomer is used as a therapeutic agent for treatment of APC and axin-related disease states.

### Assay Systems

The invention provides assay systems and screening methods for identifying specific modulators of MAPCAX activity. As used herein, an "assay system" encompasses all the components required for performing and analyzing results of an assay that detects and/or measures a particular event. In general, primary assays are used to identify or confirm a modulator's specific biochemical or molecular effect with respect to the MAPCAX nucleic acid or protein. In general, secondary assays further assess the activity of a MAPCAX modulating agent identified by a primary assay and may confirm

that the modulating agent affects MAPCAX in a manner relevant to the APC and axin pathways. In some cases, MAPCAX modulators will be directly tested in a secondary assay.

- In a preferred embodiment, the screening method comprises contacting a suitable assay system comprising a MAPCAX polypeptide or nucleic acid with a candidate agent under conditions whereby, but for the presence of the agent, the system provides a reference activity (e.g. binding activity), which is based on the particular molecular event the screening method detects. A statistically significant difference between the agent-biased activity and the reference activity indicates that the candidate agent modulates MAPCAX activity, and hence the APC and axin pathways. The MAPCAX polypeptide or nucleic acid used in the assay may comprise any of the nucleic acids or polypeptides described above.

### **Primary Assays**

- The type of modulator tested generally determines the type of primary assay.

#### *Primary assays for small molecule modulators*

- For small molecule modulators, screening assays are used to identify candidate modulators. Screening assays may be cell-based or may use a cell-free system that recreates or retains the relevant biochemical reaction of the target protein (reviewed in Sittampalam GS *et al.*, Curr Opin Chem Biol (1997) 1:384-91 and accompanying references). As used herein the term "cell-based" refers to assays using live cells, dead cells, or a particular cellular fraction, such as a membrane, endoplasmic reticulum, or mitochondrial fraction. The term "cell free" encompasses assays using substantially purified protein (either endogenous or recombinantly produced), partially purified or crude cellular extracts. Screening assays may detect a variety of molecular events, including protein-DNA interactions, protein-protein interactions (e.g., receptor-ligand binding), transcriptional activity (e.g., using a reporter gene), enzymatic activity (e.g., via a property of the substrate), activity of second messengers, immunogenicity and changes in cellular morphology or other cellular characteristics. Appropriate screening assays may use a wide range of detection methods including fluorescent, radioactive, colorimetric, spectrophotometric, and amperometric methods, to provide a read-out for the particular molecular event detected.

- Cell-based screening assays usually require systems for recombinant expression of MAPCAX and any auxiliary proteins demanded by the particular assay. Appropriate methods for generating recombinant proteins produce sufficient quantities of proteins that retain their relevant biological activities and are of sufficient purity to optimize activity
- 5 and assure assay reproducibility. Yeast two-hybrid and variant screens, and mass spectrometry provide preferred methods for determining protein-protein interactions and elucidation of protein complexes. In certain applications, when MAPCAX-interacting proteins are used in screens to identify small molecule modulators, the binding specificity of the interacting protein to the MAPCAX protein may be assayed by various known
- 10 methods such as substrate processing (e.g. ability of the candidate MAPCAX-specific binding agents to function as negative effectors in MAPCAX-expressing cells), binding equilibrium constants (usually at least about  $10^7 \text{ M}^{-1}$ , preferably at least about  $10^8 \text{ M}^{-1}$ , more preferably at least about  $10^9 \text{ M}^{-1}$ ), and immunogenicity (e.g. ability to elicit MAPCAX specific antibody in a heterologous host such as a mouse, rat, goat or rabbit).
- 15 For enzymes and receptors, binding may be assayed by, respectively, substrate and ligand processing.

The screening assay may measure a candidate agent's ability to specifically bind to or modulate activity of a MAPCAX polypeptide, a fusion protein thereof, or to cells or membranes bearing the polypeptide or fusion protein. The MAPCAX polypeptide can be 20 full length or a fragment thereof that retains functional MAPCAX activity. The MAPCAX polypeptide may be fused to another polypeptide, such as a peptide tag for detection or anchoring, or to another tag. The MAPCAX polypeptide is preferably human MAPCAX, or is an ortholog or derivative thereof as described above. In a preferred embodiment, the screening assay detects candidate agent-based modulation of MAPCAX 25 interaction with a binding target, such as an endogenous or exogenous protein or other substrate that has MAPCAX -specific binding activity, and can be used to assess normal MAPCAX gene function.

Suitable assay formats that may be adapted to screen for MAPCAX modulators are known in the art. Preferred screening assays are high throughput or ultra high throughput 30 and thus provide automated, cost-effective means of screening compound libraries for lead compounds (Fernandes PB, Curr Opin Chem Biol (1998) 2:597-603; Sundberg SA, Curr Opin Biotechnol 2000, 11:47-53). In one preferred embodiment, screening assays uses fluorescence technologies, including fluorescence polarization, time-resolved fluorescence, and fluorescence resonance energy transfer. These systems offer means to

monitor protein-protein or DNA-protein interactions in which the intensity of the signal emitted from dye-labeled molecules depends upon their interactions with partner molecules (e.g., Selvin PR, Nat Struct Biol (2000) 7:730-4; Fernandes PB, *supra*; Hertzberg RP and Pope AJ, Curr Opin Chem Biol (2000) 4:445-451).

5 A variety of suitable assay systems may be used to identify candidate MAPCAX and APC and axin pathways modulators (e.g. U.S. Pat. Nos. 5,550,019 and 6,133,437 (apoptosis assays); U.S. Pat. No. 6,114,132 (phosphatase and protease assays), U.S. Pat. Nos. 5,976,782, 6,225,118 and 6,444,434 (angiogenesis assays), among others). Specific preferred assays are described in more detail below.

10 Protein phosphatases catalyze the removal of a gamma phosphate from a serine, threonine or tyrosine residue in a protein substrate. Since phosphatases act in opposition to kinases, appropriate assays measure the same parameters as kinase assays. In one example, the dephosphorylation of a fluorescently labeled peptide substrate allows trypsin cleavage of the substrate, which in turn renders the cleaved substrate significantly more 15 fluorescent (Nishikata M *et al.*, Biochem J (1999) 343:35-391). In another example, fluorescence polarization (FP), a solution-based, homogeneous technique requiring no immobilization or separation of reaction components, is used to develop high throughput screening (HTS) assays for protein phosphatases. This assay uses direct binding of the phosphatase with the target, and increasing concentrations of target- phosphatase increase 20 the rate of dephosphorylation, leading to a change in polarization (Parker GJ *et al.*, (2000) J Biomol Screen 5:77-88).

Glycosyltransferases mediate changes in glycosylation patterns that, in turn, may affect the function of glycoproteins and/or glycolipids and, further downstream, processes of development, differentiation, transformation and cell-cell recognition. An assay for 25 glycosyltransferase uses scintillation methods to measure the transfer of carbohydrate from radiolabeled sugar-nucleotide donor to a synthetic glycopolymers acceptor that is coupled to polyacrylamide and coated on plastic microtiter plates (Donovan RS *et al.*, Glycoconj J (1999) 16:607-615).

Assays for ATPase activity are well-known in the art, such as described in 30 Blackburn et al (Blackburn CL, *et al.*, (1999) J Org Chem 64:5565-5570). The ATPase assay is performed using the EnzCheck ATPase kit (Molecular Probes). The assays are performed using an Ultraspec spectrophotometer (Pharmacia), and the progress of the reaction are monitored by absorbance increase at 360 nm. Microtubules (1.7 mM final), kinesin ( 0.11 mM final), inhibitor (or DMSO blank at 5% final), and the EnzCheck

components (purine nucleotide phosphorylase and MESG substrate) are premixed in the cuvette in a reaction buffer (40 mM PIPES pH 6.8, 5 mM paclitaxel, 1 mM EGTA, 5 mM MgCl<sub>2</sub>). The reaction is initiated by addition of MgATP (1 mM final).

- 5         **Apoptosis assays.** Assays for apoptosis may be performed by terminal deoxynucleotidyl transferase-mediated digoxigenin-11-dUTP nick end labeling (TUNEL) assay. The TUNEL assay is used to measure nuclear DNA fragmentation characteristic of apoptosis ( Lazebnik *et al.*, 1994, Nature 371, 346), by following the incorporation of fluorescein-dUTP (Yonehara *et al.*, 1989, J. Exp. Med. 169, 1747). Apoptosis may further 10 be assayed by acridine orange staining of tissue culture cells (Lucas, R., et al., 1998, Blood 15:4730-41). Other cell-based apoptosis assays include the caspase-3/7 assay and the cell death nucleosome ELISA assay. The caspase 3/7 assay is based on the activation of the caspase cleavage activity as part of a cascade of events that occur during programmed cell death in many apoptotic pathways. In the caspase 3/7 assay (commercially available Apo- 15 ONE™ Homogeneous Caspase-3/7 assay from Promega, cat# 67790), lysis buffer and caspase substrate are mixed and added to cells. The caspase substrate becomes fluorescent when cleaved by active caspase 3/7. The nucleosome ELISA assay is a general cell death assay known to those skilled in the art, and available commercially (Roche, Cat# 1774425). This assay is a quantitative sandwich-enzyme-immunoassay which uses 20 monoclonal antibodies directed against DNA and histones respectively, thus specifically determining amount of mono- and oligonucleosomes in the cytoplasmic fraction of cell lysates. Mono and oligonucleosomes are enriched in the cytoplasm during apoptosis due to the fact that DNA fragmentation occurs several hours before the plasma membrane breaks down, allowing for accumulation in the cytoplasm. Nucleosomes are not present in 25 the cytoplasmic fraction of cells that are not undergoing apoptosis. An apoptosis assay system may comprise a cell that expresses a MAPCAX, and that optionally has defective APC and axin function (e.g. APC and axin is over-expressed or under-expressed relative to wild-type cells). A test agent can be added to the apoptosis assay system and changes in induction of apoptosis relative to controls where no test agent is added, identify 30 candidate APC and axin modulating agents. In some embodiments of the invention, an apoptosis assay may be used as a secondary assay to test a candidate APC and axin modulating agents that is initially identified using a cell-free assay system. An apoptosis assay may also be used to test whether MAPCAX function plays a direct role in apoptosis. For example, an apoptosis assay may be performed on cells that over- or under-express

MAPCAX relative to wild type cells. Differences in apoptotic response compared to wild type cells suggests that the MAPCAX plays a direct role in the apoptotic response. Apoptosis assays are described further in US Pat. No. 6,133,437.

5       **Cell proliferation and cell cycle assays.** Cell proliferation may be assayed via bromodeoxyuridine (BRDU) incorporation. This assay identifies a cell population undergoing DNA synthesis by incorporation of BRDU into newly-synthesized DNA. Newly-synthesized DNA may then be detected using an anti-BRDU antibody (Hoshino *et al.*, 1986, Int. J. Cancer 38, 369; Campana *et al.*, 1988, J. Immunol. Meth. 107, 79), or by  
10 other means.

Cell proliferation is also assayed via phospho-histone H3 staining, which identifies a cell population undergoing mitosis by phosphorylation of histone H3. Phosphorylation of histone H3 at serine 10 is detected using an antibody specific to the phosphorylated form of the serine 10 residue of histone H3. (Chadlee,D.N. 1995, J. Biol. Chem 270:20098-  
15 105). Cell Proliferation may also be examined using [<sup>3</sup>H]-thymidine incorporation (Chen, J., 1996, Oncogene 13:1395-403; Jeoung, J., 1995, J. Biol. Chem. 270:18367-73). This assay allows for quantitative characterization of S-phase DNA syntheses. In this assay, cells synthesizing DNA will incorporate [<sup>3</sup>H]-thymidine into newly synthesized DNA. Incorporation can then be measured by standard techniques such as by counting of  
20 radioisotope in a scintillation counter (e.g., Beckman LS 3800 Liquid Scintillation Counter). Another proliferation assay uses the dye Alamar Blue (available from Biosource International), which fluoresces when reduced in living cells and provides an indirect measurement of cell number (Voigt-Harbin SL *et al.*, 1998, In Vitro Cell Dev Biol Anim 34:239-46). Yet another proliferation assay, the MTS assay, is based on in  
25 vitro cytotoxicity assessment of industrial chemicals, and uses the soluble tetrazolium salt, MTS. MTS assays are commercially available, for example, the Promega CellTiter 96® AQueous Non-Radioactive Cell Proliferation Assay (Cat.# G5421).

Cell proliferation may also be assayed by colony formation in soft agar (Sambrook et al., Molecular Cloning, Cold Spring Harbor (1989)). For example, cells transformed  
30 with MAPCAX are seeded in soft agar plates, and colonies are measured and counted after two weeks incubation.

Cell proliferation may also be assayed by measuring ATP levels as indicator of metabolically active cells. Such assays are commercially available, for example Cell Titer-Glo™, which is a luminescent homogeneous assay available from Promega.

Involvement of a gene in the cell cycle may be assayed by flow cytometry (Gray JW et al. (1986) Int J Radiat Biol Relat Stud Phys Chem Med 49:237-55). Cells transfected with a MAPCAX may be stained with propidium iodide and evaluated in a flow cytometer (available from Becton Dickinson), which indicates accumulation of cells 5 in different stages of the cell cycle.

Accordingly, a cell proliferation or cell cycle assay system may comprise a cell that expresses a MAPCAX, and that optionally has defective APC and axin function (e.g. APC and axin is over-expressed or under-expressed relative to wild-type cells). A test agent can be added to the assay system and changes in cell proliferation or cell cycle 10 relative to controls where no test agent is added, identify candidate APC and axin modulating agents. In some embodiments of the invention, the cell proliferation or cell cycle assay may be used as a secondary assay to test a candidate APC and axin modulating agents that is initially identified using another assay system such as a cell-free assay system. A cell proliferation assay may also be used to test whether MAPCAX function 15 plays a direct role in cell proliferation or cell cycle. For example, a cell proliferation or cell cycle assay may be performed on cells that over- or under-express MAPCAX relative to wild type cells. Differences in proliferation or cell cycle compared to wild type cells suggests that the MAPCAX plays a direct role in cell proliferation or cell cycle.

20         **Angiogenesis.** Angiogenesis may be assayed using various human endothelial cell systems, such as umbilical vein, coronary artery, or dermal cells. Suitable assays include Alamar Blue based assays (available from Biosource International) to measure proliferation; migration assays using fluorescent molecules, such as the use of Becton Dickinson Falcon HTS FluoroBlock cell culture inserts to measure migration of cells 25 through membranes in presence or absence of angiogenesis enhancer or suppressors; and tubule formation assays based on the formation of tubular structures by endothelial cells on Matrigel® (Becton Dickinson). Accordingly, an angiogenesis assay system may comprise a cell that expresses a MAPCAX, and that optionally has defective APC and axin function (e.g. APC and axin is over-expressed or under-expressed relative to wild-type cells). A test agent can be added to the angiogenesis assay system and changes in 30 angiogenesis relative to controls where no test agent is added, identify candidate APC and axin modulating agents. In some embodiments of the invention, the angiogenesis assay may be used as a secondary assay to test a candidate APC and axin modulating agents that is initially identified using another assay system. An angiogenesis assay may also be used

to test whether MAPCAX function plays a direct role in cell proliferation. For example, an angiogenesis assay may be performed on cells that over- or under-express MAPCAX relative to wild type cells. Differences in angiogenesis compared to wild type cells suggests that the MAPCAX plays a direct role in angiogenesis. U.S. Pat. Nos. 5,976,782, 5 6,225,118 and 6,444,434, among others, describe various angiogenesis assays.

**Hypoxic induction.** The alpha subunit of the transcription factor, hypoxia inducible factor-1 (HIF-1), is upregulated in tumor cells following exposure to hypoxia in vitro. Under hypoxic conditions, HIF-1 stimulates the expression of genes known to be 10 important in tumour cell survival, such as those encoding glyolytic enzymes and VEGF. Induction of such genes by hypoxic conditions may be assayed by growing cells transfected with MAPCAX in hypoxic conditions (such as with 0.1% O<sub>2</sub>, 5% CO<sub>2</sub>, and balance N<sub>2</sub>, generated in a Napco 7001 incubator (Precision Scientific)) and normoxic conditions, followed by assessment of gene activity or expression by Taqman®. For 15 example, a hypoxic induction assay system may comprise a cell that expresses a MAPCAX, and that optionally has defective APC and axin function (e.g. APC and axin is over-expressed or under-expressed relative to wild-type cells). A test agent can be added to the hypoxic induction assay system and changes in hypoxic response relative to controls where no test agent is added, identify candidate APC and axin modulating agents. In some 20 embodiments of the invention, the hypoxic induction assay may be used as a secondary assay to test a candidate APC and axin modulating agents that is initially identified using another assay system. A hypoxic induction assay may also be used to test whether MAPCAX function plays a direct role in the hypoxic response. For example, a hypoxic induction assay may be performed on cells that over- or under-express MAPCAX relative 25 to wild type cells. Differences in hypoxic response compared to wild type cells suggests that the MAPCAX plays a direct role in hypoxic induction.

**Cell adhesion.** Cell adhesion assays measure adhesion of cells to purified adhesion proteins, or adhesion of cells to each other, in presence or absence of candidate 30 modulating agents. Cell-protein adhesion assays measure the ability of agents to modulate the adhesion of cells to purified proteins. For example, recombinant proteins are produced, diluted to 2.5g/mL in PBS, and used to coat the wells of a microtiter plate. The wells used for negative control are not coated. Coated wells are then washed, blocked with 1% BSA, and washed again. Compounds are diluted to 2x final test concentration

and added to the blocked, coated wells. Cells are then added to the wells, and the unbound cells are washed off. Retained cells are labeled directly on the plate by adding a membrane-permeable fluorescent dye, such as calcein-AM, and the signal is quantified in a fluorescent microplate reader.

5 Cell-cell adhesion assays measure the ability of agents to modulate binding of cell adhesion proteins with their native ligands. These assays use cells that naturally or recombinantly express the adhesion protein of choice. In an exemplary assay, cells expressing the cell adhesion protein are plated in wells of a multiwell plate. Cells expressing the ligand are labeled with a membrane-permeable fluorescent dye, such as  
10 BCECF, and allowed to adhere to the monolayers in the presence of candidate agents. Unbound cells are washed off, and bound cells are detected using a fluorescence plate reader.

15 High-throughput cell adhesion assays have also been described. In one such assay, small molecule ligands and peptides are bound to the surface of microscope slides using a microarray spotter, intact cells are then contacted with the slides, and unbound cells are washed off. In this assay, not only the binding specificity of the peptides and modulators against cell lines are determined, but also the functional cell signaling of attached cells using immunofluorescence techniques *in situ* on the microchip is measured (Falsey JR et al., Bioconjug Chem. 2001 May-Jun;12(3):346-53).

20 **Tubulogenesis.** Tubulogenesis assays monitor the ability of cultured cells, generally endothelial cells, to form tubular structures on a matrix substrate, which generally simulates the environment of the extracellular matrix. Exemplary substrates include Matrigel™ (Becton Dickinson), an extract of basement membrane proteins containing laminin, collagen IV, and heparin sulfate proteoglycan, which is liquid at 4°C and forms a solid gel at 37°C. Other suitable matrices comprise extracellular components such as collagen, fibronectin, and/or fibrin. Cells are stimulated with a pro-angiogenic stimulant, and their ability to form tubules is detected by imaging. Tubules can generally be detected after an overnight incubation with stimuli, but longer or shorter time frames  
25 may also be used. Tube formation assays are well known in the art (e.g., Jones MK et al., 1999, Nature Medicine 5:1418-1423). These assays have traditionally involved stimulation with serum or with the growth factors FGF or VEGF. Serum represents an undefined source of growth factors. In a preferred embodiment, the assay is performed with cells cultured in serum free medium, in order to control which process or pathway a  
30

candidate agent modulates. Moreover, we have found that different target genes respond differently to stimulation with different pro-angiogenic agents, including inflammatory angiogenic factors such as TNF-alpa. Thus, in a further preferred embodiment, a tubulogenesis assay system comprises testing a MAPCAX's response to a variety of factors, such as FGF, VEGF, phorbol myristate acetate (PMA), TNF-alpha, ephrin, etc.

**Cell Migration.** An invasion/migration assay (also called a migration assay) tests the ability of cells to overcome a physical barrier and to migrate towards pro-angiogenic signals. Migration assays are known in the art (e.g., Paik JH et al., 2001, J Biol Chem 10 276:11830-11837). In a typical experimental set-up, cultured endothelial cells are seeded onto a matrix-coated porous lamina, with pore sizes generally smaller than typical cell size. The matrix generally simulates the environment of the extracellular matrix, as described above. The lamina is typically a membrane, such as the transwell polycarbonate membrane (Corning Costar Corporation, Cambridge, MA), and is generally part of an upper chamber that is in fluid contact with a lower chamber containing pro-angiogenic stimuli. Migration is generally assayed after an overnight incubation with stimuli, but longer or shorter time frames may also be used. Migration is assessed as the number of cells that crossed the lamina, and may be detected by staining cells with hematoxylin solution (VWR Scientific, South San Francisco, CA), or by any other method for determining cell number. In another exemplary set up, cells are fluorescently labeled and migration is detected using fluorescent readings, for instance using the Falcon HTS FluoroBlok (Becton Dickinson). While some migration is observed in the absence of stimulus, migration is greatly increased in response to pro-angiogenic factors. As described above, a preferred assay system for migration/invasion assays comprises testing a MAPCAX's response to a variety of pro-angiogenic factors, including tumor angiogenic and inflammatory angiogenic agents, and culturing the cells in serum free medium.

**Sprouting assay.** A sprouting assay is a three-dimensional *in vitro* angiogenesis assay that uses a cell-number defined spheroid aggregation of endothelial cells ("spheroid"), embedded in a collagen gel-based matrix. The spheroid can serve as a starting point for the sprouting of capillary-like structures by invasion into the extracellular matrix (termed "cell sprouting") and the subsequent formation of complex anastomosing networks (Korff and Augustin, 1999, J Cell Sci 112:3249-58). In an exemplary experimental set-up, spheroids are prepared by pipetting 400 human umbilical

vein endothelial cells into individual wells of a nonadhesive 96-well plates to allow overnight spheroidal aggregation (Korff and Augustin: J Cell Biol 143: 1341-52, 1998). Spheroids are harvested and seeded in 900 $\mu$ l of methocel-collagen solution and pipetted into individual wells of a 24 well plate to allow collagen gel polymerization. Test agents 5 are added after 30 min by pipetting 100  $\mu$ l of 10-fold concentrated working dilution of the test substances on top of the gel. Plates are incubated at 37°C for 24h. Dishes are fixed at the end of the experimental incubation period by addition of paraformaldehyde. Sprouting intensity of endothelial cells can be quantitated by an automated image analysis system to determine the cumulative sprout length per spheroid.

10

#### *Primary assays for antibody modulators*

For antibody modulators, appropriate primary assays test is a binding assay that tests the antibody's affinity to and specificity for the MAPCAX protein. Methods for testing antibody affinity and specificity are well known in the art (Harlow and Lane, 1988, 15 1999, *supra*). The enzyme-linked immunosorbant assay (ELISA) is a preferred method for detecting MAPCAX-specific antibodies; others include FACS assays, radioimmunoassays, and fluorescent assays.

In some cases, screening assays described for small molecule modulators may also be used to test antibody modulators.

20

#### *Primary assays for nucleic acid modulators*

For nucleic acid modulators, primary assays may test the ability of the nucleic acid modulator to inhibit or enhance MAPCAX gene expression, preferably mRNA expression. In general, expression analysis comprises comparing MAPCAX expression in like 25 populations of cells (*e.g.*, two pools of cells that endogenously or recombinantly express MAPCAX) in the presence and absence of the nucleic acid modulator. Methods for analyzing mRNA and protein expression are well known in the art. For instance, Northern blotting, slot blotting, ribonuclease protection, quantitative RT-PCR (*e.g.*, using the TaqMan®, PE Applied Biosystems), or microarray analysis may be used to confirm that 30 MAPCAX mRNA expression is reduced in cells treated with the nucleic acid modulator (*e.g.*, Current Protocols in Molecular Biology (1994) Ausubel FM *et al.*, eds., John Wiley & Sons, Inc., chapter 4; Freeman WM *et al.*, Biotechniques (1999) 26:112-125; Kallioniemi OP, Ann Med 2001, 33:142-147; Blohm DH and Guiseppi-Elie, A Curr Opin Biotechnol 2001, 12:41-47). Protein expression may also be monitored. Proteins are most

commonly detected with specific antibodies or antisera directed against either the MAPCAX protein or specific peptides. A variety of means including Western blotting, ELISA, or in situ detection, are available (Harlow E and Lane D, 1988 and 1999, *supra*).

- In some cases, screening assays described for small molecule modulators,  
5 particularly in assay systems that involve MAPCAX mRNA expression, may also be used to test nucleic acid modulators.

### **Secondary Assays**

- Secondary assays may be used to further assess the activity of MAPCAX-modulating agent identified by any of the above methods to confirm that the modulating agent affects MAPCAX in a manner relevant to the APC and axin pathways. As used herein, MAPCAX-modulating agents encompass candidate clinical compounds or other agents derived from previously identified modulating agent. Secondary assays can also be used to test the activity of a modulating agent on a particular genetic or biochemical pathway or to test the specificity of the modulating agent's interaction with MAPCAX.  
10  
15

- Secondary assays generally compare like populations of cells or animals (e.g., two pools of cells or animals that endogenously or recombinantly express MAPCAX) in the presence and absence of the candidate modulator. In general, such assays test whether treatment of cells or animals with a candidate MAPCAX-modulating agent results in changes in the APC and axin pathways in comparison to untreated (or mock- or placebo-treated) cells or animals. Certain assays use "sensitized genetic backgrounds", which, as used herein, describe cells or animals engineered for altered expression of genes in the APC and axin or interacting pathways.  
20

25           ***Cell-based assays***

- Cell based assays may detect endogenous APC and axin pathways activity or may rely on recombinant expression of APC and axin pathways components. Any of the aforementioned assays may be used in this cell-based format. Candidate modulators are typically added to the cell media but may also be injected into cells or delivered by any other efficacious means.  
30

### ***Animal Assays***

A variety of non-human animal models of normal or defective APC and axin pathways may be used to test candidate MAPCAX modulators. Models for defective APC

and axin pathways typically use genetically modified animals that have been engineered to mis-express (e.g., over-express or lack expression in) genes involved in the APC and axin pathways. Assays generally require systemic delivery of the candidate modulators, such as by oral administration, injection, etc.

5 In a preferred embodiment, APC and axin pathways activity is assessed by monitoring neovascularization and angiogenesis. Animal models with defective and normal APC and axin are used to test the candidate modulator's affect on MAPCAX in Matrigel® assays. Matrigel® is an extract of basement membrane proteins, and is composed primarily of laminin, collagen IV, and heparin sulfate proteoglycan. It is provided as a sterile liquid at 4°C, but rapidly forms a solid gel at 37°C. Liquid  
10 Matrigel® is mixed with various angiogenic agents, such as bFGF and VEGF, or with human tumor cells which over-express the MAPCAX. The mixture is then injected subcutaneously(SC) into female athymic nude mice (Taconic, Germantown, NY) to support an intense vascular response. Mice with Matrigel® pellets may be dosed via oral  
15 (PO), intraperitoneal (IP), or intravenous (IV) routes with the candidate modulator. Mice are euthanized 5 - 12 days post-injection, and the Matrigel® pellet is harvested for hemoglobin analysis (Sigma plasma hemoglobin kit). Hemoglobin content of the gel is found to correlate the degree of neovascularization in the gel.

In another preferred embodiment, the effect of the candidate modulator on  
20 MAPCAX is assessed via tumorigenicity assays. Tumor xenograft assays are known in the art (see, e.g., Ogawa K et al., 2000, Oncogene 19:6043-6052). Xenografts are typically implanted SC into female athymic mice, 6-7 week old, as single cell suspensions either from a pre-existing tumor or from *in vitro* culture. The tumors which express the MAPCAX endogenously are injected in the flank,  $1 \times 10^5$  to  $1 \times 10^7$  cells per mouse in a  
25 volume of 100 µL using a 27gauge needle. Mice are then ear tagged and tumors are measured twice weekly. Candidate modulator treatment is initiated on the day the mean tumor weight reaches 100 mg. Candidate modulator is delivered IV, SC, IP, or PO by bolus administration. Depending upon the pharmacokinetics of each unique candidate modulator, dosing can be performed multiple times per day. The tumor weight is assessed  
30 by measuring perpendicular diameters with a caliper and calculated by multiplying the measurements of diameters in two dimensions. At the end of the experiment, the excised tumors maybe utilized for biomarker identification or further analyses. For immunohistochemistry staining, xenograft tumors are fixed in 4% paraformaldehyde,

0.1M phosphate, pH 7.2, for 6 hours at 4°C, immersed in 30% sucrose in PBS, and rapidly frozen in isopentane cooled with liquid nitrogen.

In another preferred embodiment, tumorigenicity is monitored using a hollow fiber assay, which is described in U.S. Pat No. US 5,698,413. Briefly, the method comprises  
5 implanting into a laboratory animal a biocompatible, semi-permeable encapsulation device containing target cells, treating the laboratory animal with a candidate modulating agent, and evaluating the target cells for reaction to the candidate modulator. Implanted cells are generally human cells from a pre-existing tumor or a tumor cell line. After an appropriate period of time, generally around six days, the implanted samples are harvested for  
10 evaluation of the candidate modulator. Tumorigenicity and modulator efficacy may be evaluated by assaying the quantity of viable cells present in the macrocapsule, which can be determined by tests known in the art, for example, MTT dye conversion assay, neutral red dye uptake, trypan blue staining, viable cell counts, the number of colonies formed in soft agar, the capacity of the cells to recover and replicate in vitro, etc.

15 In another preferred embodiment, a tumorigenicity assay use a transgenic animal, usually a mouse, carrying a dominant oncogene or tumor suppressor gene knockout under the control of tissue specific regulatory sequences; these assays are generally referred to as transgenic tumor assays. In a preferred application, tumor development in the transgenic model is well characterized or is controlled. In an exemplary model, the "RIP1-Tag2"  
20 transgene, comprising the SV40 large T-antigen oncogene under control of the insulin gene regulatory regions is expressed in pancreatic beta cells and results in islet cell carcinomas (Hanahan D, 1985, Nature 315:115-122; Parangi S et al, 1996, Proc Natl Acad Sci USA 93: 2002-2007; Bergers G et al, 1999, Science 284:808-812). An "angiogenic switch," occurs at approximately five weeks, as normally quiescent capillaries in a subset  
25 of hyperproliferative islets become angiogenic. The RIP1-TAG2 mice die by age 14 weeks. Candidate modulators may be administered at a variety of stages, including just prior to the angiogenic switch (e.g., for a model of tumor prevention), during the growth of small tumors (e.g., for a model of intervention), or during the growth of large and/or invasive tumors (e.g., for a model of regression). Tumorigenicity and modulator efficacy  
30 can be evaluating life-span extension and/or tumor characteristics, including number of tumors, tumor size, tumor morphology, vessel density, apoptotic index, etc.

### Diagnostic and therapeutic uses

Specific MAPCAX-modulating agents are useful in a variety of diagnostic and therapeutic applications where disease or disease prognosis is related to defects in the APC and axin pathways, such as angiogenic, apoptotic, or cell proliferation disorders.

- 5 Accordingly, the invention also provides methods for modulating the APC and axin pathways in a cell, preferably a cell pre-determined to have defective or impaired APC and axin function (e.g. due to overexpression, underexpression, or misexpression of APC and axin, or due to gene mutations), comprising the step of administering an agent to the cell that specifically modulates MAPCAX activity. Preferably, the modulating agent produces
- 10 a detectable phenotypic change in the cell indicating that the APC and axin function is restored. The phrase "function is restored", and equivalents, as used herein, means that the desired phenotype is achieved, or is brought closer to normal compared to untreated cells. For example, with restored APC and axin function, cell proliferation and/or progression through cell cycle may normalize, or be brought closer to normal relative to
- 15 untreated cells. The invention also provides methods for treating disorders or disease associated with impaired APC and axin function by administering a therapeutically effective amount of a MAPCAX -modulating agent that modulates the APC and axin pathways. The invention further provides methods for modulating MAPCAX function in a cell, preferably a cell pre-determined to have defective or impaired MAPCAX function,
- 20 by administering a MAPCAX -modulating agent. Additionally, the invention provides a method for treating disorders or disease associated with impaired MAPCAX function by administering a therapeutically effective amount of a MAPCAX -modulating agent.

The discovery that MAPCAX is implicated in APC and axin pathways provides for a variety of methods that can be employed for the diagnostic and prognostic evaluation of diseases and disorders involving defects in the APC and axin pathways and for the identification of subjects having a predisposition to such diseases and disorders.

Various expression analysis methods can be used to diagnose whether MAPCAX expression occurs in a particular sample, including Northern blotting, slot blotting, ribonuclease protection, quantitative RT-PCR, and microarray analysis. (e.g., Current Protocols in Molecular Biology (1994) Ausubel FM *et al.*, eds., John Wiley & Sons, Inc., chapter 4; Freeman WM *et al.*, Biotechniques (1999) 26:112-125; Kallioniemi OP, Ann Med 2001, 33:142-147; Blohm and Guiseppi-Elie, Curr Opin Biotechnol 2001, 12:41-47). Tissues having a disease or disorder implicating defective APC and axin signaling that express a MAPCAX, are identified as amenable to treatment with a MAPCAX modulating

agent. In a preferred application, the APC and axin defective tissue overexpresses a MAPCAX relative to normal tissue. For example, a Northern blot analysis of mRNA from tumor and normal cell lines, or from tumor and matching normal tissue samples from the same patient, using full or partial MAPCAX cDNA sequences as probes, can

5 determine whether particular tumors express or overexpress MAPCAX. Alternatively, the TaqMan® is used for quantitative RT-PCR analysis of MAPCAX expression in cell lines, normal tissues and tumor samples (PE Applied Biosystems).

Various other diagnostic methods may be performed, for example, utilizing reagents such as the MAPCAX oligonucleotides, and antibodies directed against a

10 MAPCAX, as described above for: (1) the detection of the presence of MAPCAX gene mutations, or the detection of either over- or under-expression of MAPCAX mRNA relative to the non-disorder state; (2) the detection of either an over- or an under-abundance of MAPCAX gene product relative to the non-disorder state; and (3) the detection of perturbations or abnormalities in the signal transduction pathway mediated by

15 MAPCAX.

Thus, in a specific embodiment, the invention is drawn to a method for diagnosing a disease or disorder in a patient that is associated with alterations in MAPCAX expression, the method comprising: a) obtaining a biological sample from the patient; b) contacting the sample with a probe for MAPCAX expression; c) comparing results from step (b) with a control; and d) determining whether step (c) indicates a likelihood of the disease or disorder. Preferably, the disease is cancer. The probe may be either DNA or protein, including an antibody.

## EXAMPLES

25 The following experimental section and examples are offered by way of illustration and not by way of limitation.

### I. C. elegans Axin/APC Suppressor Screen

We have discovered that while RNAi of *apr-1* in a wildtype background does not

30 produce a Muv phenotype, *apr-1* inactivation enhances the penetrance of the Muv phenotype of the *pry-1* mutant to 95% (see also Gleason et al., *supra*). This enhancement of the *pry-1* Muv phenotype requires wildtype *bar-1/beta-catenin* and *pop-1/TCF* activity, suggesting that *apr-1* normally negatively regulates beta-catenin. beta-catenin-specific suppressor genes, when inactivated, likely suppress beta-catenin's inappropriate

transcriptional activation of target genes and, therefore, may be relevant for cancer therapy.

We designed a genetic screen to identify genes in addition to *bar-1*/beta-catenin and *pop-1*/TCF that act positively in beta-catenin signaling and, when inactivated, 5 suppress the Muv mutant phenotype of *pry-1* (*mu38*); *apr-1* (*RNAi*). The function of individual genes was inactivated by *RNAi* in *pry-1* mutant L1 larvae, in combination with *apr-1* *RNAi*, and suppression of the Muv phenotype was scored as a statistically significant increase in the proportion of adults that did not display the Muv phenotype. Suppressor genes were subsequently counterscreened to eliminate those that appeared to 10 suppress the *pry-1* (*mu38*); *apr-1* (*RNAi*) mutant non-specifically, rather than those that specifically function in beta-catenin signaling. Suppressor genes that passed two specificity assays were considered to be beta-catenin-specific suppressors. First, these suppressors, like *bar-1*/beta-catenin, do not suppress the Muv phenotype of three mutations in genes unrelated to beta-catenin signaling (*let-60/Ras*, *lin-12/Notch*, and *lin-15*). Second, these suppressors are not generally defective in the *RNAi* response, as 15 determined by co-*RNAi* with genes unrelated to beta-catenin signaling.

## II. Analysis of Table 1

BLAST analysis (Altschul et al., *supra*) was employed to identify orthologs of *C. elegans* modifiers. The columns “MAPCAX symbol”, and “MAPCAX name aliases” 20 provide a symbol and the known name abbreviations for the Targets, where available, from Genbank. “MAPCAX RefSeq\_NA or GI\_NA”, “MAPCAX GI\_AA”, “MAPCAX NAME”, and “MAPCAX Description” provide the reference DNA sequences for the MAPCAXs as available from National Center for Biology Information (NCBI), MAPCAX 25 protein Genbank identifier number (GI#), MAPCAX name, and MAPCAX description, all available from Genbank, respectively. The length of each amino acid is in the “MAPCAX Protein Length” column.

Names and Protein sequences of *C. elegans* modifiers of APC and axin from screen (Example I), are represented in the “Modifier Name” and “Modifier GI\_AA” 30 column by GI#, respectively.

Table1

MAPCA X symbol	MAPCAX name aliases	MAPCAX RefSeq_N A or GI_NA	NA SE	MAPCA X Q ID or NO RefSeq_ AA	AA SE	MAPCAX name	MAPCAX description	MAP CAX prote in lengt h	Modifi er name	Modifi er GI_AA
LOC2561 29	LOC256129   similar to UDP- GlcNAc:betaGa l beta-1,3-N- acetylglucosami nyltransferase 1; beta-1,3-N- acetylglucosami nyltransferase; beta-1,3-N- acetylglucosami nyltransferase 1; UDP- Gal:betaGlcNA c beta 1,3- galactosyltransf erase, polypeptide 6   na	XM_17195 5.1	1	22051164	27	similar to UDP- GlcNAc:betaGa l beta-1,3-N- acetylglucosami nyltransferase 1; beta-1,3-N- acetylglucosami nyltransferase; beta-1,3-N- acetylglucosami nyltransferase 1; UDP- Gal:betaGlcNA c beta 1,3- galactosyltransf erase, polypeptide 6	na	397	T15D6 .5	175090 57
MGC4655	MGC4655   hypothetical protein MGC4655	NM_03330 9.1	2	15208631	28	hypothetical protein MGC4655	transferase; transferase; UDP-galactose beta-N- acetylglucosa mine beta-1,3- galactosyltrans ferase	377	T15D6 .5	175090 57
B3GNT4	B3GNT4   B3GN-T4   beta3Gn-T4   beta-1,3-N- acetylglucosami nyltransferase bGn-T4   UDP- GlcNAc:betaGa l beta-1,3-N- acetylglucosami nyltransferase 4	NM_03076 5.1 XM_03 9199.1	3	13540527	29	UDP- GlcNAc:betaGa l beta-1,3-N- acetylglucosami nyltransferase 4	transferase, transferring glycosyl groups; acetylglucosa minyltransfера <sup>s</sup> e	378	T15D6 .5	175090 57

B3GNT3	B3GNT3   TMEM3   B3GN-T3   B3GNT-3   HP10328   B3GAL-T8   beta3Gn-T3   transmembrane protein 3   putative type II membrane protein   beta-1,3-N-acetylglucosaminylyltransferase bGnT-3   UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 3	NM_014254 6.2	7657172	30	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 3	acetylglucosaminylyltransferase	372	T15D6 .5	175090 57
B3GNT7	B3GNT7   hypothetical gene supported by AK000770   UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 7	NM_145235 6.1 XM_048735.1	21687139	31	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 7	transferase; UDP-galactose beta-N-acetylglucosamine beta-1,3-galactosyltransferase	401	T15D6 .5	175090 57
B3GNT1	B3GNT1   B3GNT   B3GN-T1   B3GN-T2   B3GNT-2   BETA3GNT   beta3gal-T5 gene   beta-1,3-N-acetylglucosaminylyltransferase bGnT-1   beta-1,3-N-acetylglucosaminylyltransferase bGnT-2   UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 1	NM_006576 7.3 NM_033252.1	9845238	32	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminylyltransferase 1	acetylglucosaminylyltransferase; N-acetyllactosamine synthase	397	T15D6 .5	175090 57
IMAGE:4907098	IMAGE:4907098   B3Gn-T6   beta-1,3-N-acetylglucosaminylyltransferase protein	NM_138707 6.1 XM_166247.1	20162576	33	beta-1,3-N-acetylglucosaminylyltransferase protein	acetylglucosaminylyltransferase	384	T15D6 .5	175090 57

CHL1	CHL1   CALL   L1CAM2   cell adhesion molecule with homology to L1CAM (close homologue of L1)   cell adhesion molecule with homology to L1CAM (close homolog of L1)	NM_00661 4.1	8	5729767	34	cell adhesion molecule with homology to L1CAM (close homolog of L1)	cell adhesion molecule	1224	lad-1	175387 00
L1CAM	L1CAM   HSAS   MASA   MIC5   SPG1   CAML1   CD171   HSAS1   N-CAML1   L1 cell adhesion molecule   neural cell adhesion molecule L1   L1 cell adhesion molecule (hydrocephalus, stenosis of aqueduct of Sylvius 1, MASA (mental retardation, aphasia, shuffling gait and adducted thumbs) syndrome, spastic paraplegia 1)	NM_00042 5.2   NM_02 4003.1	9	13435353	35	L1 cell adhesion molecule (hydrocephalus, stenosis of aqueduct of Sylvius 1, MASA (mental retardation, aphasia, shuffling gait and adducted thumbs) syndrome, spastic paraplegia 1)	cell adhesion molecule; cell adhesion molecule; cell adhesion molecule; cytoskeletal protein binding; integrin binding	1253	lad-1	175387 00
NFASC	NFASC   KIAA0756   neurofascin	XM_04680 8.8	10	27478636	36	neurofascin	cell adhesion molecule; cell adhesion molecule; protein binding; cytoskeletal protein binding; transmembrane receptor	1066	lad-1	175387 00

HUS1	HUS1   HUS1 (S. pombe) checkpoint homolog   HUS1 checkpoint homolog (S. pombe)	NM_00450 7.1	11	4758576	37	HUS1 checkpoint homolog (S. pombe)	DNA binding; protein binding; ATP binding; DNA clamp loader	280	hus-1	175079 89
HUS1b	HUS1b   similar to HUS1 checkpoint homolog (S. pombe); HUS1 (S. pombe) checkpoint homolog	NM_14895 9.1	12	22507374	38	similar to HUS1 checkpoint homolog (S. pombe); HUS1 (S. pombe) checkpoint homolog	protein binding	278	hus-1	175079 89
FLJ12735	FLJ12735   hypothetical protein FLJ12735	NM_02485 7.3	13	26080431	39	hypothetical protein FLJ12735	ATP binding; DNA clamp loader	1844	rfc-1	175632 26
RFC1	RFC1   A1   RFC   PO-GA   RECC1   MHCBFB   RFC140   replication factor C1   MHC binding factor, beta   replication factor C (activator 1) 1 (145kD)   replication factor C (activator 1) 1, 145kDa	NM_00291 3.2 NM_006081	14	15011931	40	replication factor C (activator 1) 1, 145kDa	DNA dependent adenosinetriphosphatase; enzyme activator; enzyme activator; ATP binding; DNA clamp loader	1148	rfc-1	175632 26
PPP4C	PPP4C   PP4   PPX   Protein phosphatase 4, catalytic subunit   protein phosphatase 4 (formerly X), catalytic subunit	NM_00272 0	15	4506027	41	protein phosphatase 4 (formerly X), catalytic subunit	protein serine/threonine phosphatase; protein phosphatase; protein phosphatase	307	pph-4.2	175543 98
YME1L1	YME1L1   FTSH   MEG4   YME1L   ATP-dependent metalloprotease FtsH1 homolog   YME1-like 1 (S. cerevisiae)	NM_01426 3 NM_139 312 NM_1 39313	16	21327685	42	YME1-like 1 (S. cerevisiae)	ATPase; ATPase; ATP binding; ATP binding; metallopeptidase; metallopeptidase; chaperone; peptidase; zinc binding	773	3L509	175542 64

EHD1	EHD1   PAST   HPAST   H-PAST   testilin   EH domain containing 1   homolog of Drosophila past   EH-domain containing 1	NM_006795	17	5803009	43	EH-domain containing 1	protein binding; insulin-like growth factor receptor binding	534	rme-1	175651 30
EHD2	EHD2   EH domain containing 2   EH-domain containing 2	NM_014601	18	21361462	44	EH-domain containing 2	nucleotide binding; protein binding	543	rme-1	175651 30
EHD3	EHD3   EH domain containing 3   EH-domain containing 3	NM_014600	19	7657056	45	EH-domain containing 3	nucleotide binding	535	rme-1	175651 30
EHD4	EHD4   EH domain containing 4   ortholog of rat pincher   EH-domain containing 4	NM_139265	20	21264315	46	EH-domain containing 4	collagen binding; nucleotide binding; calcium ion binding	541	rme-1	175651 30
KIAA0963	KIAA0963   FLJ00173   KIAA0963 protein	NM_014963	21	7662410	47	KIAA0963 protein	na	1366	nsh-1	175530 78
MOP3	MOP3   FLJ10701   FLJ10833   MOP-3	NM_018183	22	11990420	48	MOP-3	na	1392	nsh-1	175530 78
TNKS	TNKS   TIN1   PARPL   TINF1   TNKS1   TANKYRASE   tankyrase, TRF1-interacting ankyrin-related ADP-ribose polymerase	NM_003747	23	4507613	49	takyrase, TRF1-interacting ankyrin-related ADP-ribose polymerase	NAD+ ADP-ribosyltransferase; protein binding	1327	Ce_pm e-5	251460 18
TNKS2	TNKS2   TNKL   TANK2   tankyrase 2   tankyrase, TRF1-interacting ankyrin-related ADP-ribose polymerase 2	NM_025235	24	13376842	50	takyrase, TRF1-interacting ankyrin-related ADP-ribose polymerase 2	NAD+ ADP-ribosyltransferase; protein binding	1166	Ce_pm e-5	251460 18

KIAA043 3	KIAA0433   KIAA0433 protein	NM_01521 6	25	7662118	51	KIAA0433 protein	establishment and/or maintenance of cell polarity	1243	1G205	251441 24
KIAA037 7	KIAA0377   KIAA0377 gene product	NM_01465 9	26	7662084	52	KIAA0377 gene product	establishment and/or maintenance of cell polarity	1406	1G205	251441 24
NRCAM	NRCAM   KIAA0343   Bravo   neuronal cell adhesion molecule	NM_00501 0.1	53	4826864	54	neuronal cell adhesion molecule	cell adhesion molecule	1180	Lad-1	175387 00

### III. High-Throughput In Vitro Fluorescence Polarization Assay

Fluorescently-labeled MAPCAX peptide/substrate are added to each well of a 96-well microtiter plate, along with a test agent in a test buffer (10 mM HEPES, 10 mM NaCl, 6 mM magnesium chloride, pH 7.6). Changes in fluorescence polarization, determined by using a Fluorolite FPM-2 Fluorescence Polarization Microtiter System (Dynatech Laboratories, Inc), relative to control values indicates the test compound is a candidate modifier of MAPCAX activity.

### IV. High-Throughput In Vitro Binding Assay.

<sup>33</sup>P-labeled MAPCAX peptide is added in an assay buffer (100 mM KCl, 20 mM HEPES pH 7.6, 1 mM MgCl<sub>2</sub>, 1% glycerol, 0.5% NP-40, 50 mM beta-mercaptoethanol, 1 mg/ml BSA, cocktail of protease inhibitors) along with a test agent to the wells of a Neutralite-avidin coated assay plate and incubated at 25°C for 1 hour. Biotinylated substrate is then added to each well and incubated for 1 hour. Reactions are stopped by washing with PBS, and counted in a scintillation counter. Test agents that cause a difference in activity relative to control without test agent are identified as candidate APC and axin modulating agents.

### V. Immunoprecipitations and Immunoblotting

For coprecipitation of transfected proteins, 3 × 10<sup>6</sup> appropriate recombinant cells containing the MAPCAX proteins are plated on 10-cm dishes and transfected on the following day with expression constructs. The total amount of DNA is kept constant in each transfection by adding empty vector. After 24 h, cells are collected, washed once with phosphate-buffered saline and lysed for 20 min on ice in 1 ml of lysis buffer

containing 50 mM Hepes, pH 7.9, 250 mM NaCl, 20 mM -glycerophosphate, 1 mM sodium orthovanadate, 5 mM p-nitrophenyl phosphate, 2 mM dithiothreitol, protease inhibitors (complete, Roche Molecular Biochemicals), and 1% Nonidet P-40. Cellular debris is removed by centrifugation twice at 15,000 × g for 15 min. The cell lysate is  
5 incubated with 25 µl of M2 beads (Sigma) for 2 h at 4 °C with gentle rocking.

After extensive washing with lysis buffer, proteins bound to the beads are solubilized by boiling in SDS sample buffer, fractionated by SDS-polyacrylamide gel electrophoresis, transferred to polyvinylidene difluoride membrane and blotted with the indicated antibodies. The reactive bands are visualized with horseradish peroxidase  
10 coupled to the appropriate secondary antibodies and the enhanced chemiluminescence (ECL) Western blotting detection system (Amersham Pharmacia Biotech).

## VI. Expression analysis

All cell lines used in the following experiments are NCI (National Cancer Institute)  
15 lines, and are available from ATCC (American Type Culture Collection, Manassas, VA 20110-2209). Normal and tumor tissues are obtained from Impath, UC Davis, Clontech, Stratagene, Ardais, Genome Collaborative, and Ambion.

TaqMan® analysis is used to assess expression levels of the disclosed genes in various samples.

20 RNA is extracted from each tissue sample using Qiagen (Valencia, CA) RNeasy kits, following manufacturer's protocols, to a final concentration of 50ng/µl. Single stranded cDNA is then synthesized by reverse transcribing the RNA samples using random hexamers and 500ng of total RNA per reaction, following protocol 4304965 of Applied Biosystems (Foster City, CA).

25 Primers for expression analysis using TaqMan® assay (Applied Biosystems, Foster City, CA) are prepared according to the TaqMan® protocols, and the following criteria: a) primer pairs are designed to span introns to eliminate genomic contamination, and b) each primer pair produced only one product. Expression analysis is performed using a 7900HT instrument.

30 TaqMan® reactions are carried out following manufacturer's protocols, in 25 µl total volume for 96-well plates and 10 µl total volume for 384-well plates, using 300nM primer and 250 nM probe, and approximately 25ng of cDNA. The standard curve for result analysis is prepared using a universal pool of human cDNA samples, which is a mixture of cDNAs from a wide variety of tissues so that the chance that a target will be

present in appreciable amounts is good. The raw data are normalized using 18S rRNA (universally expressed in all tissues and cells).

For each expression analysis, tumor tissue samples are compared with matched normal tissues from the same patient. A gene is considered overexpressed in a tumor 5 when the level of expression of the gene is 2 fold or higher in the tumor compared with its matched normal sample. In cases where normal tissue is not available, a universal pool of cDNA samples is used instead. In these cases, a gene is considered overexpressed in a tumor sample when the difference of expression levels between a tumor sample and the average of all normal samples from the same tissue type is greater than 2 times the 10 standard deviation of all normal samples (i.e., Tumor – average(all normal samples) > 2 x STDEV(all normal samples)).

A modulator identified by an assay described herein can be further validated for therapeutic effect by administration to a tumor in which the gene is overexpressed. A decrease in tumor growth confirms therapeutic utility of the modulator. Prior to treating a 15 patient with the modulator, the likelihood that the patient will respond to treatment can be diagnosed by obtaining a tumor sample from the patient, and assaying for expression of the gene targeted by the modulator. The expression data for the gene(s) can also be used as a diagnostic marker for disease progression. The assay can be performed by expression analysis as described above, by antibody directed to the gene target, or by any other 20 available detection method.

**WHAT IS CLAIMED IS:**

1. A method of identifying a candidate APC and axin pathways modulating agent, said method comprising the steps of:
  - 5       (a) providing an assay system comprising a MAPCAX polypeptide or nucleic acid;
  - (b) contacting the assay system with a test agent under conditions whereby, but for the presence of the test agent, the system provides a reference activity; and
  - (c) detecting a test agent-biased activity of the assay system, wherein a difference between the test agent-biased activity and the reference activity identifies the test agent as
- 10      a candidate APC and axin pathways modulating agent.
2. The method of Claim 1 wherein the assay system comprises cultured cells that express the MAPCAX polypeptide.
- 15      3. The method of Claim 2 wherein the cultured cells additionally have defective APC and axin function.
- 20      4. The method of Claim 1 wherein the assay system includes a screening assay comprising a MAPCAX polypeptide, and the candidate test agent is a small molecule modulator.
- 25      5. The method of Claim 4 wherein the assay is a binding assay.
- 30      6. The method of Claim 1 wherein the assay system is selected from the group consisting of an apoptosis assay system, a cell proliferation assay system, an angiogenesis assay system, and a hypoxic induction assay system.
- 35      7. The method of Claim 1 wherein the assay system includes a binding assay comprising a MAPCAX polypeptide and the candidate test agent is an antibody.
- 40      8. The method of Claim 1 wherein the assay system includes an expression assay comprising a MAPCAX nucleic acid and the candidate test agent is a nucleic acid modulator.

9. The method of Claim 8 wherein the nucleic acid modulator is an antisense oligomer.

10. The method of Claim 8 wherein the nucleic acid modulator is a PMO.

5 11. The method of Claim 1 additionally comprising:

(d) administering the candidate APC and axin pathways modulating agent identified in (c) to a model system comprising cells defective in APC and axin function and, detecting a phenotypic change in the model system that indicates that the APC and axin function is restored.

10

12. The method of Claim 11 wherein the model system is a mouse model with defective APC and axin function.

15 13. A method for modulating a APC and axin pathways of a cell comprising contacting a cell defective in APC and axin function with a candidate modulator that specifically binds to a MAPCAX polypeptide, whereby APC and axin function is restored.

20 14. The method of Claim 13 wherein the candidate modulator is administered to a vertebrate animal predetermined to have a disease or disorder resulting from a defect in APC and axin function.

15. The method of Claim 13 wherein the candidate modulator is selected from the group consisting of an antibody and a small molecule.

25 16. The method of Claim 1, comprising the additional steps of:

(e) providing a secondary assay system comprising cultured cells or a non-human animal expressing MAPCAX ,

30 (f) contacting the secondary assay system with the test agent of (b) or an agent derived therefrom under conditions whereby, but for the presence of the test agent or agent derived therefrom, the system provides a reference activity; and

(g) detecting an agent-biased activity of the second assay system,  
wherein a difference between the agent-biased activity and the reference activity of the second assay system confirms the test agent or agent derived therefrom as a candidate APC and axin pathways modulating agent,

and wherein the second assay detects an agent-biased change in the APC and axin pathways.

17. The method of Claim 16 wherein the secondary assay system comprises cultured  
5 cells.
18. The method of Claim 16 wherein the secondary assay system comprises a non-human animal.
- 10 19. The method of Claim 18 wherein the non-human animal mis-expresses a APC and axin pathways gene.
- 15 20. A method of modulating APC and axin pathways in a mammalian cell comprising contacting the cell with an agent that specifically binds a MAPCAX polypeptide or nucleic acid.
21. The method of Claim 20 wherein the agent is administered to a mammalian animal predetermined to have a pathology associated with the APC and axin pathways.
- 20 22. The method of Claim 20 wherein the agent is a small molecule modulator, a nucleic acid modulator, or an antibody.
23. A method for diagnosing a disease in a patient comprising:
  - (a) obtaining a biological sample from the patient;
  - 25 (b) contacting the sample with a probe for MAPCAX expression;
  - (c) comparing results from step (b) with a control;
  - (d) determining whether step (c) indicates a likelihood of disease.
24. The method of Claim 23 wherein said disease is cancer.

## SEQUENCE LISTING

<110> EXELIXIS, INC.

<120> MAPCAXs AS MODIFIERS OF THE APC AND AXIN PATHWAYS AND METHODS OF USE

<130> EX04-003C-PC

<150> US60/443,484  
<151> 2003-01-29

<150> US60/447,358  
<151> 2003-02-11

<150> US60/461,789  
<151> 2003-04-10

<150> US60/470,684  
<151> 2003-05-14

<150> US60/479,650  
<151> 2003-06-19

<160> 54

<170> PatentIn version 3.2

<210> 1  
<211> 1194  
<212> DNA  
<213> Homo sapiens

<400> 1  
atgcgctgcc ccaagtgcct tctctgcctg tcagcactgc tcacactcct gggcctcaaa 60  
gtgtacatcg agtggacatc cgagtcccg ctcagcaagg cctaccccg ccctcggggc 120  
accccgccaa gccccacgcc agccaacctt gagcccaccc tacctgccaa cctctccacc 180  
cgccctgggcc agactatccc gctgccctt gcttactgga accagcagca gtggcggctg 240  
gggtccctgc ccagtggga cagcactgaa acgggggct gccaggctt gggggccgccc 300  
gccgccaccc agatccctga cttcgcctcc tacccttcaagg acctccggcg cttttgctg 360  
tcagcagcct gccggagctt cccacagtgg ctgcctggag gtggtgccag ccaagtctcc 420  
agctgctcag atactgatgt cccctacctg ctgttggccg tcaagtcaaga accagggcgc 480  
tttgcagaac gacaggccgt gagagagacg tggggcagtc cagctccagg gatccggctg 540  
ctcttcctgc tagggctcctcc ggtgggtgag gcggggcctg accttagactc actagtggcc 600  
tgggagagcc gtcgctacag tgacctgctg ctctggact tcctcgacgt cccattcaac 660  
cagacgctca aagacctgct gctgctggcc tggctggcc gccactgccc caccgtgagt 720  
tttgtcttgc gagctcagga cgatgcctt gtacacaccc ctgcccgtt ggctcacctg 780  
cgggccctgc cacctgcctc ggcccgaaac ctctacctgg gtgaggtctt taccctaggcc 840

atgcctctcc ggaagccagg aggacccttc tatgtccccg agtccttctt cgaagggtggc	900
tacccagcct atgcaagcgg gggtggtac gtcattgccg ggccctggc accctggctg	960
ctgcggcggt cagccgtgt ggcacccttc cccttgagg acgtctacac tggctttgc	1020
atccgagccc tgggccttgtt gccccaggcc cacccaggct tcctcacagc ctggccagca	1080
gaccgcactg cgaccactg tgcttccgc aacctgctgc tggtaacggcc cctggggccc	1140
caggccagca ttccggctctg gaaacaactg caagacccaa ggctccagtg ctga	1194

<210> 2  
 <211> 2712  
 <212> DNA  
 <213> Homo sapiens

<400> 2	
caccggtccc ctttcctgcc aacgctgcat ttggctcggtt cccggtccat ggccggccct	60
cggaccctgc gctgagccccc ggaggccagg gcgtccgggg ctgcgccact tccgagggcc	120
gagcgctgcc ggtcccgcg gtgcgacacg gccgggagga ggagaacaac gcaaggggct	180
caaccgtcgg tcgctggagc cccccccggg gcgtggcctc ccggccctc agctggggag	240
ggcggggctc gctccccct gctgccact gcgaccctta caggggaggg agggcgcagg	300
ccgcgcggag atgaggagga ggctgcgcct acgcaggac gcattgctca cgctgctcct	360
tggcgctcc ctgggcctct tactctatgc gcagcgcgac ggcgcggccc cgacggcggag	420
cgcgcgcgca gggcgaggga gggcggcacc gaggcccacc cccggacccc gcgcgttcca	480
gttacccgac gcgggtgcag ccccgccggc ctacgaaggg gacacacccgg cgccgcccac	540
gcctacggga ccctttgac ttgcggcgct atttgcgcgc caaggaccag cggcggtttc	600
cactgctcat taaccagccg cacaagtgcc gcggcgacgg cgcacccggt ggccgcccgg	660
acctgcttat tgctgtcaag tcgggtggcag aggacttcga gcggcgccaa gccgtgcgcc	720
agacgtgggg cgcggagggt cgcgtgcagg gggcgctggt gcgcgcgtg ttcttgctgg	780
gcgtgcccag gggcgcaggc tcgggccccggg ccgacgaagt tggggagggc gcgcgaaccc	840
actggcgccgc cctgctgcgg gccgagagcc ttgcgtatgc ggacatcctg ctctggccct	900
tcgacgacac ctttttaac ctaacgctca aggagatcca ctttctagcc tggccctcag	960
ctttctgccc cgacgtgcgc ttgcgtttta agggcgacgc agatgtgttc gtgaacgtgg	1020
gaaatctcct ggagttcctg gcgcgcggg accccggcgca agacctgctt gctggtgacg	1080
taatttgtca tgccggccc atccgcacgc gggctagcaa gtactacatc cccgaggccg	1140
tgtacggcct gccccctat ccggcctacg cggcgccgg tggcttgc tttccgggg	1200
ccacgctgca ccgcctggct ggccctgtg cgcaaggcga gctttcccc atcgacgacg	1260
tctttctggg catgtgtctg cagcgccctgc ggctcacgccc cgagcctcac cctgccttcc	1320

gcacctttgg catccccag cttcagccg cgccgcattt gaggcaccc tc gacccctgt	1380
tttaccgtga gctgggtgta gtgcacgggc tctcgccgc tgacatctgg cttatgtggc	1440
gcctgctgca cgggccat gggccagcct gtgcgcattc acaggctgtc gtcaggcc	1500
ccttccaatg ggactcctag ctccccacta cagccccaa agccctaaactc agacccagaa	1560
tggagccggt ttcccagatt attgccgtgt atgtggttct tccctgatca ccaggtgcct	1620
gtctccacag gatcccagg gatgggggtt aagcttggtc cctggcggtc caccctgt	1680
gaaccagttg aaacccgtgt aatggtgacc cttttagcga gccaaggctg ggtggtagat	1740
gaccatctct tgtccaacag gtcccagagc agtggatatg tctggtcctc ctagtagcac	1800
agaggtgtgt tctgggtgtgg tggcagggac ttaggaaatc ctaccactct gctggatttg	1860
gaacccctta ggctgacgca gacgtatgca gaggctctca aggccaggcc ccacagggag	1920
gtggaggggc tccggccgcc acagcctgaa ttcatgaacc tggcaggcac tttgccatag	1980
ctcatctgaa aacagatatt atgcttccca caacctctcc tgggcccagg tggctgtag	2040
caccagggat ggagccacac ataaggaca aatgagtgcg cggccttacc tagtctttcc	2100
tcacccctcg aactcacaca acaatgccag tctccactg gaggctgtat cccctcagag	2160
gagccaagga atgtcttccc ctgagatgcc accactatta atttcccat atgcttcaac	2220
caccccttg ctcaaaaaac caataccac acttaccta atacaaacat cccagcaaca	2280
gcacatggca ggccattgct gagggcacag gtgcatttatt ggagagggga tggggcagg	2340
ggataaggaa gttccccca ttccaggagg atggaaacag tcctggctgc ccctgacagt	2400
ggggatatgc aaggggctct ggccaggcca cagtccaaat gggaaagacac cagtcagtca	2460
caaaagtcgg gagcgcacaca caaacctggc tataaggccc aggaaccata taggagcctg	2520
agacaggtcc cctgcacatt catcattaaa ctatacagga tgaggctgta catgagttaa	2580
ttacaaaaga gtcatattta caaaaatctg tacacacatt tgaaaaactc acaaaattgt	2640
catctatgta tcacaagttt cttagacaaa atattaaaaa tggataaaaa ttataaaaaa	2700
aaaaaaaaaa aa	2712

<210> 3  
 <211> 1296  
 <212> DNA  
 <213> Homo sapiens

<400> 3 cacagcctga gactcatctc gttcgaccc cgccggcc gcccggccc ggcattctga	60
gcacggagac agtctccagc tgccgttcat gttccctccc cagccttctg cagccacca	120
ggaaaggggc ggttaggagtg gccttttacc aaagggaccc gcgatgtct gcaggctgt	180

ctggctggtc tcgtacagct tggctgtgct gttgctcggc tgccctgctct tcctgaggaa	240
ggcggccaag cccgcaggag acccccacggc ccaccagcct ttctgggctc ccccaacacc	300
ccgtcacagc cggtgtccac ccaaccacac agtgtcttagc gcctctctgt ccctgcctag	360
ccgtcaccgt ctcttcttga cctatcgta ctgccgaaat ttctcttatct tgctggagcc	420
ttcaggctgt tccaaggata ccttcttgcct cctggccatc aagtacacagc ctggtcacgt	480
ggagcgacgt gcggtatcc gcagcacgtg gggcagggtg gggggatggg cttagggccg	540
gcagctgaag ctggtgttcc tcctaggggt ggcaggatcc gctccccag cccagctgct	600
ggcctatgag agtagggagt ttgatgacat cctccagtgg gacttcactg aggacttctt	660
caacctgacg ctcaaggagc tgcacctgca gcgctgggtg gtggctgcct gccccaggc	720
ccatttcatg ctaaaggag atgacgatgt ctttgtccac gtccccaaacg tgttagagtt	780
cctggatggc tgggacccag cccaggacct cctgggtggaa gatgtcatcc gccaaagccct	840
gcccaacagg aacactaagg tcaaataactt catcccaccc tcaatgtaca gggccaccca	900
ctacccaccc tatgctggtg ggggaggata tgtcatgtcc agagccacag tgccgcgcct	960
ccaggctatc atggaagatg ctgaactt cccattgtat gatgtctttg tggatgttg	1020
cctgaggagg ctgggctga gccctatgca ccatgctggc ttcaagacat ttggaatccg	1080
gccccccctg gacccttag acccctgcct gtataggggg ctcctgtgg ttcaccgcct	1140
cagccccctc gagatgtgga ccatgtggc actggtgaca gatgaggggc tcaagtgtgc	1200
agctggcccc atacccacgc gctgaagggt gggttggca acagcctgag agtggactca	1260
gtgttatttc tctatcgta tgcgaaattt atgcct	1296

<210> 4  
 <211> 2195  
 <212> DNA  
 <213> Homo sapiens

<400> 4	
actctttctt cggctcgca gctgagagga gcaggttagag gggcagagggc gggactgtcg	60
tctggggag ccgcccagga ggctcctcag gccgacccca gaccctggct ggccaggatg	120
aagtatctcc ggcacccggcg gcccaatgcc accctcattc tggccatcg cgcttcacc	180
ctcctcctct tcagtctgct agtgtcacca cccacctgca aggtccagga gcagccacccg	240
gcgatccccg aggcctggc ctggccact ccacccaccc gcccagcccc ggccccgtgc	300
catgccaaca cctctatggt cacccacccg gacttcgcca cgcagccgca gcacgttcag	360
aacttcctcc tgtacagaca ctgcccacac tttccctgc tgcaaggacgt gccccctct	420
aagtgcgcgc agccggctt cctgctgctg gtgatcaagt ctcctccatcg caactatgtg	480
cgccgcgagc tgctgcggcg cacgtggggc cgcgagcgca aggtacgggg tttcagctg	540

cgcctcctct tcctggtggg cacagcctcc aaccgcacg aggcccgc aa	600
ggtaaccgg ctgctggagc tggaggcaca gactcacgga gacatcctgc agtgggactt ccacgactcc	660
ttcttcaacc tcacgctcaa gcaggtcctg ttcttacagt ggcaggagac aaggtgcgc	720
aacgccagct tcgtgctcaa cggggatgat gacgtcttg cacacacaga caacatggtc	780
ttctacctgc aggaccatga ccctggccgc cacctctcg tggggcaact gatccaaaac	840
gtgggccccca tccgggcttt ttggagcaag tactatgtgc cagaggtggt gactcagaat	900
gagcggtacc caccctattg tgggggtggt ggcttcttgc tgtcccgctt cacggccgct	960
gcctgcgcc gtgctgccc tgtcttggac atcttccccca ttgatgatgt cttcctgggt	1020
atgtgtctgg agcttgaggg actgaagcct gcctcccaca gcggcatccg cacgtctggc	1080
gtcgggctc catcgcaaca cctgtcctcc tttgaccctt gtttctaccg agacctgctg	1140
ctggtgacc accttcctacc ttatgagatg ctgctcatgt gggatgcgct gaaccagccc	1200
aacctcacct gcggcaatca gacacagatc tactgagtca gcatcagggt cccca	1260
gcccgcctc tgggctcctg tttccatagg aaggggcgac accttcctcc caggaagctg agacctttgt	1320
ggtctgagca taagggagtg ccagggaaagg tttgagggtt gatgagtgaa tattctggct	1380
ggcgaactcc tacacatcct tcaaaaccca cctggtaactg ttccagcatc ttccctggat	1440
ggctggagga actccagaaa atatccatct tcttttgtg gctgctaatt gcagaagtgc	1500
ctgtgctaga gttccaaactg tggatgcac cgtcccggtt gagtcaaagt cttaactccc	1560
tgtctcacc tactcacaga cgggatgcta agcagtgcac ctgcagtggc ttaatggcag	1620
ataagctccg tctgcagtcc cagggcagcc agaaactcct gtgtccacat agagctgacg	1680
tgagaaatat ctttcagccc aggagagagg ggtcctgatc ttaacccttt cctgggtctc	1740
agacaactca gaagggtggg gggataccag agaggtggtg gaataggacc gcccctcct	1800
tacttgtggg atcaaattgt gtaatggtgg aggtgtggc agaggaggaa ggcaagtgtc	1860
ctttgaaagt tgtgagagct cagagttct ggggtcctca ttaggagccc ccatccctgt	1920
gttccccaaag aattcagaga acagcactgg ggctggaatg atcttaatg ggcccaaggc	1980
caacaggcat atgcctcaact actgcctgga gaaggagag attcaggtcc tccagcagcc	2040
tccctcaccc agtatgtttt acagattacg gggggaccgg gtgagccagt gaccccctgc	2100
agcccccagc ttcaggcctc agtgtctgcc agtcaagctt cacaggcatt gtgatggggc	2160
agccttgggg aatataaaat tttgtgaaga cttgg	2195

<210> 5  
 <211> 1434  
 <212> DNA  
 <213> Homo sapiens

<400> 5	
gggttaaggc acctgggcca ccgcccctgc cggccctcc tcccgcggcc ggaaagagga	60
aagtgcggc gggggcgccgg cccggcttcc gtccccaccc gcccgcgtc ccgcggccc	120
gagccgtggc gccagagct gcgagccgct cgccctccg ccgtccggc ccggccgccc	180
atgtcgctgt ggaagaaaac cgtctaccgg agtctgtgcc tggccctggc cctgctcgtg	240
gccgtgacgg tgttccaacg cagtctcacc cctggtcagt ttctgcagga gcctccgcca	300
cccaccctgg agccacagaa ggcccagaag ccaaattggac agctggtaa ccccaacaac	360
ttctggaaga acccgaaaga tgtggctgca cccacgccc tggcctctca gggcccccag	420
gcctgggacg tgaccaccac taactgctca gccaatatca acttgaccctt ccagccctgg	480
ttccaggtcc tggagccgca gttccggcag tttcttttcc accggccactg ccgtacttc	540
cccatgctgc tgaaccaccc ggagaagtgc agggcgatg tctacctgct ggtgggtgtc	600
aagtccgtca tcacgcagca cgaccgccc gaggccatcc gccagacctg gggccgcgag	660
cggcagtccg cgggtggggg ccgaggcgcc gtgcgcaccc tttcctgct gggcacggcc	720
tccaagcagg aggagcgcac gcactaccag cagctgctgg cctacgaaga ccgcctctac	780
ggcgacatcc tgcagtgggg ctttctcgac accttcttca acctgaccctt caaggagatc	840
cacttcctca agtggctgga catctactgc cccacgtcc ctttcatttt caaaggcgac	900
gatgacgtct tcgtcaaccc caccaacctg cttagaatttc tggctgaccg gcagccacag	960
gaaaacctgt tcgtggcgta tgcctgcag cacgctcggc ccattcgcag gaaagacaac	1020
aaatactaca tcccgggggc cctgtacggc aaggccagct atccggcgtt tgcaggcgcc	1080
ggtggcttcc tcatggccgg cagcctggcc cggccctgc accatgcctg cgacaccctg	1140
gagctctacc cgatcgacga cgtctttctg ggcattgtgcc tggaggtgct gggcgtgcag	1200
cccacggccc acgaggcgtt caagactttc ggcatctccc ggaaccgcaa cagccgcatt	1260
aacaaggagc cgtgctttt ccgcgcattt ctcgtggtgc acaagctgct gcccctgag	1320
ctgctcgcca tggggggctt ggtgcacagc aatctcacct gctccgc当地 gctccagggt	1380
ctctgacccc agccggctt ctaggacagg ccagggcact tgctcctgag cccc	1434

<210> 6  
 <211> 2742  
 <212> DNA  
 <213> Homo sapiens

<400> 6	
ggcgccggca gcgtcagcag cggcaacaag tgccggagta gcagagccaa gccggagcag	60
tccctgcgcg cgcacaccgccc gggccgccc tccggggcgc cgcgcatttga gcgtgagctg	120
cgccggcgcg cgggctgagc cgcgcggagc gcccggacgt ggatgtggcc gcgtatctccc	180

gccttggccc cggcccgcc gagctggagc tgctccgga caagatatga gaaatgagtg 240  
ttggacgtcg aagaataaag ttgttggtt tcctgatgtat ggcaaatgtc ttcatttatt 300  
ttattatgg agtctccaaa agcagtagcc aagaaaaaaa tggaaaagg gaagtaataa 360  
tacccaaaga gaagttctgg aagatatcta cccctccgga ggcatactgg aaccgagagc 420  
aagagaagct gaaccggcag tacaaccca tcctgagcat gctgaccaac cagacggggg 480  
aggcgggcag gctctccaat ataagccatc tgaactactg cgaacctgac ctgaggggtca 540  
cgtcgggtt tacgggtttt aacaacttgc cggacagatt taaagacttt ctgctgtatt 600  
tgagatgccg caattattca ctgcttatag atcagccgga taagtgtgca aagaaacctt 660  
tcttgtgct ggcgattaag tccctactc cacatttgc cagaaggcaa gcaatccggg 720  
aatcctgggg ccaagaaagc aacgcaggga accaaacggt ggtgcgagtc ttccctgctgg 780  
gccagacacc cccagaggac aaccaccccg acctttcaga tatgctgaaa tttgagagtg 840  
agaagcacca agacattctt atgtgaaact acagagacac tttcttcaac ttgtctctga 900  
aggaagtgct gtttctcagg tggtaagta cttcctgccc agacactgag tttgtttca 960  
agggcgatga cgatgtttt gtgaacaccc atcacatcct gaattacttg aatagtttat 1020  
ccaagaccaa agccaaagat ctcttcatag gtgatgtgat ccacaatgct ggacctcatc 1080  
gggataagaa gctgaagtac tacatccag aagttgtta ctctggcctc taccacccct 1140  
atgcaggggg aggggggttc ctctactccg gccacctggc cctgaggctg taccatatca 1200  
ctgaccaggt ccatctctac cccattgatg acgtttatac tggaatgtgc cttcagaaac 1260  
tcggcctcgt tccagagaaa cacaaggct tcaggacatt tgatatcgag gagaaaaaca 1320  
aaaataacat ctgctcctat gtagatctga tgtagtaca tagtagaaaa cctcaagaga 1380  
tgattgatat ttggtctcaag ttgcagagtg ctcattttaa atgctaaaat agatacaaac 1440  
tcaattttgc atagaaaggt gtattttgaa tagtccccat gttgtttct cacatttagag 1500  
taatttctat attaaaccat gaaaattgcc tttatgagtg ataccattt gagggcctct 1560  
aaacccttca atttggtact cacgtgaaga gggaaagcgg aagatggtaa tttttttta 1620  
tggatgatat ggcaggatga ttggttctga tcttaccggc tagtggtcat ttttaaaaaa 1680  
cttgcacct cttatctgaa atcctgttcc tggaaatttgg ccattttaaag tgattttgtt 1740  
tgcccttttc tataatattc ctactccca taataatgac tgatttattt gttattcagg 1800  
tatttataaa cctattggct acaaagactt tgtaaactt tatccagtgg tttcgtgaa 1860  
atggaattat gtttattttt atgggatttg ggttaaatttt aaattgtcta gaaaactgaa 1920  
atttcagtttgc tcaatttttttgc aattcagttt tcaatttgc gaaatttccct gccacccaa 1980  
cagtattttt gtgtgttaat taattttgc aatgagaat catggtgtga cactcatcta 2040

atttatcttg ttgtgatgtt atggtcataa taaggagaaa gagggtttaa ttttcttgt	2100
atttggtttc ctggtggtat catagtgtaa ttttagtatt tgaaaatcag tgtgattcct	2160
taatggccaa ctgaagattt aattgccgtt aacaaccata tcgtgttagt gaattttcaa	2220
tatggaccag gaaggcatat gtattttgaa cttgagtgaa aaggttgaag ttacagactt	2280
ttgcatagat ggtttgcag tttaaaattt cagaatttgtt tattgccata tttcacatg	2340
ctgcttatac aagatttata ttgagtagta actgcttccc tgtctatgta gaagtgcctg	2400
tgtttttatt tattgttcag atcaaagacc aaaacatttt cttaaatata ttttatgtaa	2460
tattttattt gtatacagtg ttgtgatgtt aatatttaac tagagcatgtt tattttaaat	2520
gttaagggtgt aacatatgtt aaataaaaact gttttttttt aattttaaaa tttgtttttt	2580
gggggtatgtt actactagag tttaaaattt tgccaaacta ttacttataat gtactattgtt	2640
gtaacataact ttcttgagat attttgtttt atagaattgtt aggttcttat cagatgggat	2700
actggggact ataaacaatgtt gaaataaaagc cactgttattt tt	2742

<210> 7  
<211> 1155  
<212> DNA  
<213> *Homo sapiens*

<400> 7  
atggctttc cctgccgcag gtccctgact gccaagactc tggcctgcct cctggtggc 60  
gtgagtttct tagcaactgca gcagtggttc ctccaggcgac caaggcccc gcgggaggag 120  
aggccccgc aggaggagac gccagagggt cccaccgacg ctcccgccgc tgacgagccg 180  
ccctcgaggc tcgtccccgg gccccgtgc gtggcgaacg ctcggcgaa cgccacggcc 240  
gacttcgagc agctgcccgc gcgcattccag gacttcctgc ggtaccgcca ctggcccac 300  
ttcccgctgc ttgggacgc accggccaag tgcgccggcg gccgaggcgt gttcctgctc 360  
ctggcggtga agtcggcgcc tgagcaactac gagcgacgacg agctcatccg gcgcacgtgg 420  
gggcaagagc gcagctacgg cgggcccga gtgcggccgc tcttctatt gggcaccccg 480  
ggccccgagg acgaggcgcg cgccggagcgg ctggcgagc tggtggcgct ggaggcgccg 540  
gagcacggcg acgtgctgca gtgggccttc gcggacacct tcctcaacct cacgctcaag 600  
cacctgcact tgctcgactg gctggctgca cgctgcccgc acgcgcgtt tctgctcagc 660  
ggcgacgacg acgtgttcgt gcacaccgccc aacgttagtcc gttcctgca ggcgacgcca 720  
ccccggccgc acctgttctc cggccagctc atggagggtt ccgtgcccatt ccgcacagc 780  
tggagcaagt acttcgtgccc gccgcagctc ttcccccgggtt ccgttaccc ggtgtactgc 840  
agcgccggcg gcttcctcctt gtccggcccc acggccccgg ccctgcgcgc ggccgcccgc 900

cacaccccg	tcttccccat	cgacgacgcc	tacatggca	tgtgtctgga	gcgcgcggc	960
ctggcgccca	gcggccacga	gggcatccgg	cccttcggcg	tgcagctgcc	tggcgcacag	1020
cagtccctct	tcgacccttg	catgtaccgc	gagttgctgc	tagtgcacccg	cttcgcgccc	1080
tacgagatgc	tgctcatgtg	gaaggcgctg	cacagccccg	cgctcagctg	tgaccgggga	1140
caccgggtct	cctga					1155

<210> 8  
<211> 7642  
<212> DNA  
<213> Homo sapiens

<400> 8							
cggaccctgc	gcgcccccgt	ccggctccc	ggccggctcg	ggggagaagg	cgcggaggg	60	
gaggcgccgg	acagatcgcg	tttcggaggc	ggcgccagg	ctgtaaactg	caaaccataa	120	
tcctgtctta	atactgcaaa	caaatcatag	tggaactaag	gggaacttaa	tttactgttt	180	
ccaggttaac	taaggctctca	gctgtaaacc	aaaagtgaga	ggagacatta	agattttcat	240	
tcttaccggg	ttgtcttctt	cctgaagagc	aatggagccg	cttttacttg	gaagaggact	300	
aatcgatat	ctaattttcc	tcctgttaaa	attctaaaaa	gcaattgaaa	taccatcttc	360	
agttcaacag	gttccaacaa	tcataaaaca	gtcaaaagtc	caagttgcct	ttcccttcga	420	
ttagtatttt	caaattgaat	gtgaagctaa	aggaaatcca	gaaccaacat	ttcgtggac	480	
taaggatggc	aacccttttt	attcactga	ccatcgata	attccatcga	acaattcagg	540	
aacattcagg	atcccaaacg	aggggcacat	atctcacttt	caagggaaat	accgctgctt	600	
tgcttcaaatt	aaactggaa	tcgctatgtc	agaagaaata	gaatttata	ttccaagtgt	660	
tccaaaattc	ccaaaagaaa	aaattgaccc	tcttgaagtg	gaggagggag	atccaattgt	720	
cctcccatgc	aatccccc	aaggcctccc	acctttacac	atttattgga	tgaatattga	780	
attagaacac	atcgaacaag	atgaaagagt	atacatgagc	caaaagggag	atctataactt	840	
cgcaaacgtg	gaagaaaagg	acagtcgcaa	tgactactgt	tgctttgctg	catttccaag	900	
attaaggact	attgtacaga	aatgc当地	gaaactaaca	gttaacagtt	taaagcatgc	960	
taatgactca	agttcatcca	cagaaattgg	ttccaaggca	aattccatca	agcaaagaaa	1020	
acccaaactg	ctgtgcctc	ccactgagag	tggcagttag	tcttcaatta	ccatcctcaa	1080	
agggaaaatc	ttgctgcttg	agtgtttgc	tgaaggcttg	ccaaactccac	aggttgattg	1140	
gaacaaaatt	ggtggtgact	taccaaagg	gagagaagca	aaagaaaatt	atggcaagac	1200	
tttgaagata	gagaatgtct	cctaccagga	caaaggaaat	tatcgctgca	cagccagcaa	1260	
tttcttggga	acagccactc	acgattttca	cgttatagta	gaagagcctc	ctcgctggac	1320	
aaagaagcct	cagagtgctg	tgtatagcac	cggaagcaat	ggcatcttgt	tatgtgaggc	1380	

tgaaggagaa cctcaaccca caatcaagtg gagagtcaat ggctccccag ttgacaatca	1440
tccatttgct ggtgatgttgc ttcccag ggaaatcagt tttaccaacc ttcaacccaaa	1500
tcatactgct gtgtaccagt gtgaagcctc aaatgtccat ggaactatcc ttgccaatgc	1560
caatattgtat gttgtggatg tccgtccatt gatacaaacc aaagatggag aaaattacgc	1620
tacagtggtt gggtaacagtgc ctttcttaca ttgcgagttc tttgcttcac ctgaggcagt	1680
cgtgtcctgg cagaagggtgg aagaagtggaa accccotggag ggcaggcggt atcatatcta	1740
tgaaaatggc acattgcaga tcaacagaac caccgaagaa gatgctgggt cttactcatg	1800
ttgggttagaa aatgctatag gaaaaactgc agtcacagcc aatttggata ttagaaatgc	1860
tacaaaactt agagtttctc ctaagaatcc tcgtatcccc aaattgcata tgcttgaatt	1920
acattgtgaa agcaaatgtg actcacattt gaaacacagt ttgaagttgt cctggagtaa	1980
agatggagaa gccttgaaa ttaatggcac agaagatggc aggataatta ttgatggagc	2040
taatttggacc atatctaatg taactttaga ggaccaaggt atttactgct gttcagctca	2100
tactgctcta gacagtgcctg ccgatataac tcaagtaact gttcttgatg ttccggatcc	2160
accagaaaac cttcacttgt ctgaaagaca gaacaggagt gttcggctga cctgggaagc	2220
tggagctgac cacaacagca atattagcga gtatattgtt gaatttgaag gaaacaaaga	2280
agaggctgga aggtgggagg aactgaccag agtccaagga aagaaaacca cagttatctt	2340
acctttggct ccatttgcata gataccaggta cagggtcata gccgtgaacg aagtagggag	2400
aagtcagccct agccagccgt cagaccatca tgaaacacca ccagcagctc cagataggaa	2460
tccacaaaac ataagggttc aagcctctca acccaaggaa atgattataa agtggagcc	2520
tttgaatcc atggagcaga atggaccagg cctagagtac agagtgcacct ggaagccaca	2580
gggagcccca gtggagtggg aagaagaaac agtcacaaac cacacattgc gggtgatgac	2640
gcctgctgtc tatgcccattt atgatgtcaa ggtccaggct atcaatcaac taggatctgg	2700
gcctgaccct cagtcagtga ctctctattc tggagaagac tattcctgata cagctccagt	2760
gatccatggg gtggacgtta taaacagtac attagttaa gttacctggta caacagttcc	2820
aaaggacaga gtacatggac gtctgaaagg ctatcagata aattgggtggaa aaacaaaaag	2880
tctgttggat ggaagaacac atcccaaaga agtgaacatt ctaagattt caggacaaag	2940
aaactctgga atggttcattt ctttagatgc ctttagtgaa tttcattttaa cagtcattgc	3000
ctataactct aaaggagctg gtcctgaaag tgaggcttat atattcaaa caccagaagg	3060
agtacctgaa cagccaaactt ttctaaaggt catcaaagtt gataaaagaca ctgccacttt	3120
atcttgggaa ctacctaaga aattaaatgg aaacttaact ggctatctt tgcaatata	3180
gataataaat gacacccatcg agattggaga attaaatgtat attaacatccaactccatc	3240

aaagcccagc tggcacctct caaacctgaa tgcaactacc aagtacaaat tctacttgag	3300
ggcttgcact tcacagggct gtggaaaacc gatcacggag gaaagctcca ccttaggaga	3360
agggagtaaa ggtatcgga agatatcagg agtaaatctt actcaaaga ctcacccagt	3420
agaggtatt gagccggag ctgaacatat agttcgcta atgactaaga attggggcga	3480
taacgatagc attttcaag atgtaattga gacaagaggg agagaatatg ctggttata	3540
tgatgacatc tccactcaag gctggtttat tggactgatg tgtgcgattg ctctctcac	3600
actactatta ttaactgttt gctttgtgaa gaggaataga ggtggaaagt actcagttaa	3660
agaaaaggaa gatttgcattc cagacccaga aattcagtca gtaaaagatg aaacctttgg	3720
tgaatacagt gacagtgatg aaaagcctct caaaggaagc cttcggtccc ttaataggga	3780
tatgcagccct actgaaaagtg ctgacagctt agtcaatac ggagagggag accatggct	3840
cttcagtgaa gatggatcat ttattggtgc ctacgctgga tctaaggaga agggatctgt	3900
tgaaagcaat ggaagttcta cagcaacttt tccccttcgg gcataaacac aacatatgta	3960
agcaacgcta ctggttcacc ccaaccttcc atatttatct gttcaaagga gcaagaactt	4020
tcatatagga atagaaacat gctggccgaa gatttcatcc agaagtcaac atcctgcaat	4080
tatgttggaaa agagtagtac tttcttcaaaa atataaaatg ccaagcactt caggcctatg	4140
ttttgcttat attgtttca ggtgctaaa atgcaaaaca caaaacaaat cctgcattta	4200
gatacacccctc aactaaatcc aaagtccccca ttcatatat tccatatttgc cctgatttt	4260
ctattcggtg tgttgcata gatgttgcta cttgggggt ttttctccgt atgcacatttgc	4320
gtatacagtc tctgagaact ggcttggta ctggcttca ctacaggta aaagaccata	4380
agcaaactgg ttattnaaaa tgtaaaaagg aatatgaaag tcttattaaa acacttcatt	4440
gaaaatatac agtctaaatt tattattnaa atttacttag caaaagtctt aggtgaacaa	4500
tcaactagta ttgttgagc tcctatttgc ccagagatgg tcataattaa acagaagttt	4560
acgttttca gtttcaacat gaattttttt atttctgtca gttatgacat ccacgagcat	4620
cactttttgt gtctgtttt tttttttct tggactaaat tcaactgcat ggaagcggtg	4680
gtcagaaggt tgtttatac gagaacaggc agaaagtgcc cattgttcag gattctaata	4740
gctacatcta cttaatatct tcattctaa attgactgct ttaccttt tctcatgttt	4800
atataatggt atgcttgcat atatttcatg aatacattgt acatattatg ttaatattta	4860
cacaatttaa aatatagatg tgtttattt tgaagtgaga aaatgaacat taacaggcat	4920
gtttgtacag ctagaatata ttagtaagat actgttttc gtcattccag agctacaact	4980
aataacacga ggttccaaag ctgaagactt tgtataaaagt atttgggttt tggttctgtta	5040
ttgcttctt tcaacagttt caaaataaaa tatcatacaa atattgaggg aaatgttttc	5100

atattttca aaataggaaa ttattgttga atgtacatct accccagccc ctcaaaagaa	5160
aaactgttta catagaaaatt cctacacata cgtttgcgta tatgttattt taaacatctt	5220
tgtggtgaga atttttccc cgatattctc cttctgtcaa agtcagaaca aattcaggga	5280
atttattttc tggcagttgt gctccagtcc tttaaaatt gtacatgaac atgttttaga	5340
aacaatatgg aggatgatgc atacatgtcg gtcaagttca gcgcgtcgaca ttttatggaa	5400
agattttttt aacccatcca cgaaatactt aactactgtt taagtgaatt gacttatttc	5460
acttttagttt ttgaactgtg attattggta tactgttata tcctcaactt ggatttatgg	5520
taaccccttt tagttcatgg agacaaaaat ttggggatttata tataatagtc agcgcaggaa	5580
tgcacatgga atatctactt gtcctttga acctcacgag tcacccagaa tgtatagaca	5640
ggaaaagcat gtcttattta aaactgtaat ttatggctc aggatctgac cgcaagtcccg	5700
ggagtaagca tttcaaaagg ggaaggcagt gtggcccta ccctgtgtga atgtgaggat	5760
gtagacatcc atcagtgcaa ctcgagctcc atcctcctcc gatttctaag gttccagttt	5820
tctggaggga cagtcatcat gtttgattt atctggaga aaactgtggt gcacagcttg	5880
tgaggaggc aaggttgtga cgttcgagct tagttctgggt gttattctgt ctccctttct	5940
ttgtcatcag ccaaaacgtg gttttaaag agagtcatgc aggttagaaa taatgtcaaa	6000
aatattttagg aatttaataa ctttaagtc agaaaactaaa acaaatactg aaatattagc	6060
tcttcctaca cttcgtgttc cccttagct gcctgaaaat caagattgct cctactcaga	6120
tcttctgagt ggctaaaact tatggatatg aaaaatgaga ttgaatgtg actatgctt	6180
gctatcattt ttaccttcc tcaatactat ttggcaacta ctggactct tcagcacaaa	6240
aggaatagat ctatgattga ccctgatttt aattgtgaaa ttatatgatt catatatttt	6300
atgaatcaga ataaccctca aataaaataa atctaagtcg gttaaaatgg atttcatgat	6360
tttccctcag aaaatgagta acggagtccc acggcgtgca atggtaatta taaattgggt	6420
atgcttggttt gcaaattgcc cactcgtgat aagtcaacag ccaatattta aaactttgtt	6480
cgttactggc tttaccctaa ctttctctag tctactgtca atatcattt aatgttaattt	6540
attgtatata gtctcaagaa tggttgggg gcatttttttgccttgcgttca atatcatttt	6600
tggaaaatc caaattctct tcctggctcc agcactgatt ttgtacataa acattaggca	6660
ggttgcttaa cttttttatt tcaaaactctc tcaactctaa agtgctaata ataatctcag	6720
ttaccttatac tttgtcacag ggtgttctt tttatgttataaaaatggaa aatgataaaa	6780
gctaagatgc cttctaactt cataagcaaa ctttaacta attatgtatc tgaaagtac	6840
ccccacatac caactcaact ttttcctgt gaacacataa atatattttt atagaaaaac	6900
aaatctacat aaaataaaatc tactgttttag tgagcgttat gacttgcata tgccattgaa	6960

aattattaat cagaagaaaa ttaaggcaggg tctttgctat acaaaagtgt tttccactaa	7020
tttgcataat gtatttataa gaaaaatgtg aatttggtgg ttttattcta tcggtataaa	7080
ggcatcgata ttttagatgc acccggtttt gtaaaaatgt agagcacaat ggaattatgc	7140
tggaaagtctc aaataatatt ttttcctat ttatactca tggaagagat aagctaaaga	7200
ggggacaata atgagaaaatg ttgggtgtct tttctaagca tttaaaacat aattgccaat	7260
tgaaacccta aatatgttta cataccatta agatatgatt catgtaacaa tgttaaatta	7320
attataatgg gattgggttt gttatctgtg gtagtatata tcctagtgtt cctatagtga	7380
aataagttagg gttcagccaa agcttcttt gtttgtacc ttaaattgtt cgattacgtc	7440
atcaaaagag atgaaaggta ttagaaacag gttcacgtga ttacctttt ctttggctt	7500
ggattaatat tcatagtaga actttataaa acgtgttgtt attgttaggtg gtgttgtat	7560
tatgctttagt actatgtatg gtttggaaat atttcatta tacatgaaat tcaactttcc	7620
aaataaaaagt tctacttcat gt	7642

<210> 9  
 <211> 4525  
 <212> DNA  
 <213> Homo sapiens

<400> 9	
gcgcggtgcc gccgggaaag atggtcgtgg cgctgcggta cgtgtggcct ctccctct	60
gcagccccctg cctgcattatc cagatccccg aggaatatga aggacaccat gtgatggagc	120
cacctgtcat cacggAACAG tctccacggc gcctgggtgt cttccccaca gatgacatca	180
gcctcaagtgc tgaggccagt ggcaagcccg aagtgcagtt ccgctggacg agggatggtg	240
tccacttcaa acccaaggaa gagctgggtg tgaccgtgtt ccagtcgccc cactctggct	300
ccttcaccat cacggcaac aacagcaact ttgctcagag gttccagggc atctaccgct	360
gctttgccag caataagctg ggcaccccca tgtcccatga gatccggctc atggccgagg	420
gtgcccccaa gtggccaaag gagacagtga agcccggtga ggtggaggaa gggagtcag	480
tgggtctgcc ttgcaaccct ccccccaagtgc cagacccctt ccggatctac tggatgaaca	540
gcaagatctt gcacatcaag caggacgagc gggtagcgt gggccagaac ggcaacctct	600
actttgccaa tgtgttcacc tccgacaacc actcagacta catctgccac gcccacttcc	660
caggcaccag gaccatcatt cagaaggaac ccattgaccc ccgggtcaag gccaccaaca	720
gcatgattga caggaagccg cgcctgctt tccccacca ctccagcagc cacctgggtgg	780
ccttgcaggg gcagccattt gtcctggagt gcatcgccga gggcttccc acgcccacca	840
tcaaatggct ggcacccagt ggcccatgc cagccgaccg tgtcacccatc cagaaccaca	900

acaagaccct gcagctgctg aaagtggcg aggaggatga tggcgagtac cgctgcctgg	960
ccgagaactc actggcagt gcccggcatg cgtactatgt caccgtggag gctgccccgt	1020
actggctgca caagccccag agccatctat atggccagg agagactgcc cgccctggact	1080
gccaagtcca gggcaggccc caaccagagg tcacctggag aatcaacggg atccctgtgg	1140
aggagctggc caaagaccag aagtaccgga ttacagcgtgg cgccctgatc ctgagcaacg	1200
tgcagcccaag tgacacaatg gtgacccaat gtgaggcccg caaccggcac gggctttgc	1260
tggccaatgc ctacatctac gttgtccagc tgccagccaa gatcctgact gcggacaatc	1320
agacgtacat ggctgtccag ggcagcactg cctaccttct gtgcaaggcc ttcggagcgc	1380
ctgtgcccag ttttcgtgg ctggacgagg atggacaac agtgcttcag gacgaacgct	1440
tcttcccta tgccaatggg accctggca ttcgagacct ccaggccaaat gacaccggac	1500
gctacttctg cctggctgcc aatgaccaaa acaatgttac catcatggct aacctgaagg	1560
ttaaagatgc aactcagatc actcaggggc cccgcagcac aatcgagaag aaaggttcca	1620
gggtgacctt cacgtgccag gcctcctttg acccctcctt gcagcccagc atcacctggc	1680
gtggggacgg tcgagacctc caggagctt gggacagtga caagtacttc atagaggatg	1740
ggcgcttgtt catccacagc ctggactaca gcgaccaggg caactacagc tgcgtggcca	1800
gtaccgaact ggatgtggtg gagagtaggg cacagctttt ggtgggtggg agccctggc	1860
cgggtgccacg gctgggtctg tccgacctgc acctgctgac gcagagccag gtgcgcgtgt	1920
cctggagtcc tgcagaagac cacaatgccc ccattgagaa atatgacatt gaatttgagg	1980
acaaggaaat ggcgcctgaa aaatggtaca gtctggcaa gttccaggg aaccagacct	2040
ctaccaccct caagctgtcg ccctatgtcc actacacctt tagggttact gccataaaca	2100
aatatggccc cggggagccc agcccggtct ctgagactgt ggtcacaccc gaggcagccc	2160
cagagaagaa ccctgtggat gtgaaggggg aaggaaatga gaccaccaat atggtcatca	2220
cgtggaaagcc gtcgggtgg atggactgga acgccccca gttcagtgac cgctgtcagt	2280
ggccccccta ggggacacga gggccctggc aggacagat tgcagcgcac cccttcctgg	2340
tgggttccaa cacgtccacc ttctgtccct atgagatcaa agtccaggcc gtcaacagcc	2400
agggcaaggg accagagccc caggtcacta tcggctactc tggagaggac taccccccagg	2460
caatccctga gctggaaggc attgaaatcc tcaactcaag tgccgtctg gtcaagtggc	2520
ggccgggtgga cctggcccag gtcaaggccc acctccgcgg atacaatgtg acgtactgg	2580
gggagggcag tcagaggaag cacagcaaga gacatatcca caaagaccat gtgggtgtgc	2640
ccgccaacac caccagtgtc atcctcagtg gttgcggcc ctatagctcc taccacctgg	2700
agggtcaggc cttaaacggg cgaggatcg ggcccgccag cgagttcacc ttcaagcaccc	2760

cagagggagt gcctggccac cccgaggcgt tgcacctgga gtgccagtcg aacaccagcc	2820
tgctgctgcg ctggcagccc ccactcagcc acaacggcgt gtcacccgc tacgtgtct	2880
cctaccaccc cctggatgag gggggcaagg ggcaactgtc cttcaacctt cgggaccccg	2940
aacttcggac acacaacctg accgatctca gcccccacct gcggtaccgc ttccagcttc	3000
aggccaccac caaagagggc cctggtaag ccatcgtagc ggaaggaggc actatggcct	3060
tgtctggat ctcagatttt ggcaacatct cagccacagc gggtaaaaac tacagtgtcg	3120
tctcctgggt ccccaaggag gccagtgca acttcagggtt ccatacttg ttcaaagcct	3180
tggagaaga gaagggtggg gttccctt cgccacagta tgtcagctac aaccagagct	3240
cctacacgca gtgggacctg cagcctgaca ctgactacga gatccacttg tttaggaga	3300
ggatgttccg gcaccaaatg gctgtgaaga ccaatggcac aggccgcgtg aggctccctc	3360
ctgctggctt cgccactgag ggctggttca tcggcttgtt gagtgccatc atccctctgc	3420
tcctcgctt gtcatcctc tgcttcatca agcgcagcaa gggcggcaaa tactcagtga	3480
aggataagga ggacacccag gtggactctg aggcccacc gatgaaagat gagaccttcg	3540
gcgagtacag gtccctggag agtgacaacg aggagaaggc ctttggcagc agccagccat	3600
cgctcaacgg ggacatcaag cccctggca gtgacgacag cctggccgat tatggggca	3660
gcgtggatgt tcagttcaac gaggatggtt cgttcattgg ccagtacagt ggcaagaagg	3720
agaaggaggc ggcagggggc aatgacagct cagggccac ttccccatc aaccctgccc	3780
tggccctaga atagttggagt ccaggacagg agatgtgtc cccctggctt tggatccag	3840
gccctccct ctccagcagg cccatggag gctggagttg gggcagagga gaacttgctg	3900
cctcggatcc cttcttacc acccggtccc cacttattt caaaaaccca gtcacccccc	3960
ttcctggca cacgctgctc tgccccagct tggcagatc tcccacatgc cagggccctt	4020
tgggtgtgt tttgcagcc catttggca gagaggctgt gttttgggg agaagaagta	4080
ggggtggccc gaaagggtct ccgaaatgct gtctttctt ctcctgact ggggcagac	4140
atgggtgggt ctccctcagga ccagggttgg cacccccc ctccttccagc cactccccag	4200
ccagcctggc tggactggg aacagaactc ggtgtccccca ccatctgtc tctttcttt	4260
gccatctctg ctccaaacccg gatgggagcc gggcaaactg gccgcgggg caggggaggc	4320
catctggaga gcccagagtc cccccactcc cagcatcgca ctctggcagc accgccttt	4380
cccgccgccc agcccacccc atggccggct ttcaggagct ccatacacac gctgccttcg	4440
gtacccacca cacaacatcc aagtggcctc cgtcaactacc tggctgcggg gcgggcacac	4500
ctcctccac tgcccaactgg ccggc	4525



ccgcctggag	gtcaaagacc	ccaccaggat	ctaccggatg	cccgaggacc	aggtgtggccag	1800
aaggggcacc	acggtgcagc	tggagtgtcg	ggtgaagcac	gaccctccc	tgaaactcac	1860
cgtctcctgg	ctgaaggatg	acgagccgct	ctatattgga	aacaggatga	agaaggaaga	1920
cgactccctg	accatcttg	gggtggcaga	gcgggaccag	ggcagttaca	cgtgtgtcgc	1980
cagcaccgag	ctagaccaag	acctggccaa	ggcctacctc	accgtgctag	ctgatcaggc	2040
cactccaact	aaccgtttgg	ctgccctgcc	caaaggacgg	ccagaccggc	cccgggacct	2100
ggagctgacc	gacctggccg	agaggagcgt	gcggctgacc	tggatccccg	gggatgctaa	2160
caacagcccc	atcacagact	acgtcgtcca	gtttgaagaa	gaccagttcc	aacctggggt	2220
ctggcatgac	cattccaagt	accccggcag	cgttaactca	gccgtcctcc	ggctgtcccc	2280
gtatgtcaac	taccagttcc	gtgtcattgc	catcaacgag	gttgggagca	gccaccccg	2340
cctcccatcc	gagcgctacc	gaaccagtgg	agcacccccc	gagtccaatc	ctggtgacgt	2400
gaagggagag	gggaccagaa	agaacaacat	ggagatcacg	tggacgccc	tgaatgccac	2460
ctcggcctt	ggcccaacc	tgcgctacat	tgtcaagtgg	aggcggagag	agactcgaga	2520
ggcctggAAC	aacgtcacag	tgtgggctc	tcgctacgtg	gtggggcaga	ccccagtcta	2580
cgtccctat	gagatccgag	tccaggctga	aaatgacttc	gggaaggggcc	ctgagccaga	2640
gtccgtcatac	ggttactccg	gagaagattt	acccagtgcc	cctaggcggt	tccgagtccg	2700
gcagcccaac	ctggagacaa	tcaacctgga	atgggatcat	cctgagcatc	caaatgggat	2760
catgatttgg	tacactctca	aatatgtggc	ctttaacggg	accaaagttag	gaaagcagat	2820
agtggaaaac	ttctctccca	atcagaccaa	gttcacggtg	caaagaacgg	accccgtgtc	2880
acgctaccgc	tttaccctca	gcgccaggac	gcaggtgggc	tctgggaag	ccgtcacaga	2940
ggagtcacca	gcaccccccga	atgaagctac	tccaaaccgca	gcttacacca	acaaccaagc	3000
ggacatcgcc	acccagggct	ggttcattgg	gcttatgtgc	gccatcgccc	tcctgggtct	3060
gatcctgctc	atcgctgtt	tcatcaagag	gagtcgcggc	ggcaagtacc	cagtacgaga	3120
aaagaaggat	gttcccttg	gccctgaaga	ccccaaaggaa	gaggatggct	catttgacta	3180
tagtgtatgag	gacaacaagc	ccctgcaggg	cagtcagaca	tctctggacg	gcaccatcaa	3240
gcagcaggag	agtgacgaca	gcctggtgga	ctatggcgag	ggtggcgagg	gtcagttcaa	3300
tgaagacggc	tccttcatcg	gccagtacac	ggtaaaaaag	gacaaggagg	aaacagaggg	3360
caacgaaagc	tcagaggcca	cgtcacctgt	caatgctatc	tactctctgg	cctaacggag	3420
cccacccagg	cacagccacc	actttgcaag	tgggaggagg	ggagaagggg	agacaaaacc	3480
actgcagacc	taccacgaag	ccaccaccac	cttcagtaac	aagggtacga	tatgggggtc	3540
tgccaagctg	tgaggaccag	tagccaccaa	gccacccaca	agccccctcc	caatgacccc	3600

ccttcagccc cgggtgccac cagtgtggga gagctggagc cgtggctgag ctcagctgga	3660
gggagcctgg ccccttgcac ggtctcgca gCACCCGAG CGTCCACCA CACTGTCCGC	3720
ccttggcctc ggcacacgct cacctttct gttggttacg ggacttctca ttgtcttaat	3780
ttcgctttgt gcatcttccc ctccagaccc aatgtctctg ttttctttc ctttgaaaaa	3840
agagtccctg gaaaagaaaag aataagtgga ttctccccca ggagaatcct ttttgaaag	3900
ataggcaaaa aggattgaat ccataccaga acacgttagac aggctgtaat attcaaacc	3960
cctctgggcc aatgcatttc caaaggatgc ctttctcgcc atatgcctcc cctggccccc	4020
agcccctctg cctcggcctt gtcagttgct gagctggct tggctcctt ctggaaaatg	4080
acagtatttt tggcagggag aaggtgcgca ggcctcctt ctgctcttg gttgggtgg	4140
gaggtgtgtt tacctcttgc tcctcattcc tccctgcac ttttctctgg aatatctaag	4200
atgtgagctg cattgactct gaagacgtt gaggaacagg agtgggcaact gatagaaagg	4260
acttcaacgc cagtgactgt gtacctccag cagaagaaaa tcaggtgtct ggtcttgggg	4320
gcactgtgct cacttctaga gagaagaaaa aggctgggtt tggacttcat gcctcctcac	4380
tgtgaaaacc caggatgttag atagaggccc acacccaccc tgtccagagt agagggcacc	4440
tgtccacgtg gccagggccc atgctgccac cctcgaggag atgaaacct gtgatgcaaa	4500
agctttgctg gtgtgtttgg ggcataaggc actgctcccc ttccatctct agcagatatc	4560
atcttctcag ccaattctgtt aggacactga ggctttgtt tgacatggtg tgtggatgg	4620
gacttgcccc aagactgagc aggtctgccc tttcattggg gtgtatacat atcaggggac	4680
cctgaggtgg caccgtactc agtgttgtga tgccccacc tagggaggac tcaatgctct	4740
ttgttatgcct tatcagcgc tgatctgtcc ctggagttcc caggttccca gctccccccc	4800
tgcaagcctt gaagcctcca gcaacgtcta tctccaggag agcagggcag cctatgcaag	4860
tgttccggca ggaccagaag tggccactca acacacgtcc tcaccagtca ccccaaatgg	4920
agacactcgc agacctgcag ttctcagacc gaaggtgctg ctttcattctt ccccacctaa	4980
tgcatttttg aaaccaaagg tacctagcct cagagacaga gaaggagagg gggagatctt	5040
gagtctaaga ggaacccttt agctgtttct gcagctgagg tcagttcag gagggtgaag	5100
ctgatccccca ggaatggatc agctgccatg ggcacagcct ccgatttgag cctctccggc	5160
tgccagccag gggcctggg ccagaccagg gctctgtcct tccgtgactt tattaaagca	5220
acatttgcca catgtttaag ccgc	5244

<210> 11  
 <211> 2122  
 <212> DNA  
 <213> Homo sapiens

<400> 11	
ctcagggcgc gacctttct gttaccaaca gaggccgccc gcggctgcgc catccgcggc	60
atgaagtttc gggccaagat cgtggacggg gcctgtctga accacttcac acgaatcagt	120
aacatgatag ccaagcttgc caaaacctgc accctccgca tcagccctga taagcttaac	180
ttcatccttt gtgacaagct ggctaattgga ggagttagca tgtggtgtga gctggaacag	240
gagaacttct tcaacgaatt tcaaattggag ggtgtctctg cagaaaacaa tgagatttat	300
tttagagctaa catcgaaaaa cttatctgaa gccttgaaga ctgcccagaa tgccaggcgt	360
ttgaaaatca aactgactaa taaacacttt ccctgcctca cggtctccgt ggagctgtta	420
tctatgtcaa gcagtagccg cattgtgacc catgacatcc ccataaaggt gattcctagg	480
aaattgtgga aggacttaca agaaccggtg gtcccagatc ctgatgttag tatttattta	540
ccagtcttga agactatgaa gagtgttgtg gaaaaaatga aaaacatcag caatcacctt	600
gttattgaag caaacctaga tggagaattt aatttggaaaa tagaaaactga attagtatgt	660
gttacaactc attttaaaga tcttggaaat cctccattag cctctgaaag cacccatgag	720
gacagaaaacg tggAACACAT ggctgaagt cacatagata ttaggaagct cctacagttt	780
cttgctggac aacaagtaaa tcccacaaag gccttatgca atattgtgaa taacaagatg	840
gtgcattttt atctgcttca tgaagacgtg tcccttcagt atttcatccc tgcaactgtcc	900
tagcaccctg tcgctggagt tggcatgcag agactttgtc agatgggag aggccgcagg	960
tgttgtttc tgatcaactgg tctgtccct cacagcaccc cacatcgaca cactgtactt	1020
atttgtccct ctctaaccatt ttaactaaaa gttgattcaa caacacacag ttggataaac	1080
atatacattc atgttgctca tgtctgttt gctttgttt taagacactg aaaagaaaaag	1140
ctagaattta ttatttcaga cttaaagaa caatttctca ttgatgttgt gaaaatcgtc	1200
atgtatTTAG acttggtgta gtagccagaa ttctaaagc tttttttttt gagcttggta	1260
ctttccctcc aggcagaggc tctagcttag cacggctgt agcgacact cagtttgca	1320
tttcagtgtg ttccccccgc tgctcctgcc cttggagcc cagtgcacaga aagaacagcc	1380
tctgtcaccc cgccgcact gccttggta ctcagagcac tgtgggtgt cacagctgca	1440
gcatttggag tctctctt gctgaggact caagcccacc tgatccact ccccttttga	1500
tgcctagaga gctggcccag ccaacacagc tcttagctgg gagctccccc tgccattcca	1560
actagtttct tcctggggcc agttttgggt ttaggttgta attccttata tttttttttt	1620
ccacagtgttca tcggatctgt ctttctggaa agaagaccct tctattttaga gtagaaacaa	1680
acgaaaacttc taaggtatca tctgtgtttaa gtgtgagac catatttctt tgatgtttct	1740
gaacatcaaa gctgattcag tactggtaga tgtgctcatt ctccctgaaa catacccatc	1800

atattgtcct attataatta catctcattg tcctgtggag gtggacatga taaacattat 1860  
ctttgtttt cttgtttgt tttgttttag acggctcat tctgtcaccc agactggagt 1920  
gcagtgccac aatcatggct caccgcattg acctccttgg ctcaaggcat cctccacct 1980  
cagcttcctg actagctggg actactggtg tgcaccacca cacccagcta atttcaatt 2040  
tttcatagag acagggtctc actgtgttgt ccaggagttg gagaccagcc cggcaacaa 2100  
agtgagaccc cgtctctact tt 2122

<210> 12  
<211> 837  
<212> DNA  
<213> Homo sapiens

<400> 12  
atgaagtttc gcgccaagat caccggcaaa ggctgtctag agctgttcat tcacgtcagc 60  
ggcaccgtcg cgaggcttagc gaaggcttcgc gtgcctccgc tgcccccgtga cagcctgtgc 120  
ttcggccccg cgggttccgg cggcctccac gaggccaggc tgtggtgcga ggtgcggcag 180  
ggggccttcc agcagtttcg catggaaggt gtctcggaag atctcgatga gatccacctg 240  
gagctgacgg cggagcacct gtcccggcg gcgagaagcg cagcgggcgc gtctccctg 300  
aagctgcagc tgaccaccaa gcgcgcgc tccctcacgg tggcggtgga gctggtctcg 360  
tccctgggcc gcgctcgcag cgtggtgcac gatctgccc tgccgggtct tcccaggaga 420  
gtgtggcggg actgcctgccc gcccagcctg cgccgcctccg acgcgagcat ccgcctgccc 480  
cgctggagga cgctgaggag catcggtggaggatggcga acgtggcag tcacgtgctg 540  
gtggaagcaa acctcagtgg caggatgacc ctgagttatag agacggaggt ggtgtccatt 600  
caaagttatt ttaaaaatct tggaaaccct ccccaagtccg ctgtgggtgt gcctgaaaac 660  
agagacctgg agagcatggt gcaagtgcgg gtggacaatc ggaagcttct gcagttttt 720  
gagggacagc aaatacatcc tacgacggcc ctgtgcaata tttgggacaa tactttctt 780  
cagcttgcatt tggttcaaga atatgtctct cttcagtatt tcattcctgc cttgtaa 837

<210> 13  
<211> 6245  
<212> DNA  
<213> Homo sapiens

<400> 13  
tttcctccga agctctgtgg tccgatctgc ggtccgcttg ctttccctgc ccggtcccga 60  
gcgcctcagcc tgaagcgccg ctttcgaggg caccctgcac acactggccg cgccctcaggg 120  
atctcattgc ccgcgttttc tcattgcctc tttccgtgtt cgattcggt gatctggcc 180  
cagcctccgc tcccgctctc tgcgggtggg cgccggggaa tccgaaacgg ctcagcagaa 240

tcccaagcagc ttgctgctac tggagcgggc cgccctccatg gcctccaggc aggccgggct 300  
ggaccgcgtg aggtcctagg agacgggatt ccgggaagcg gggagtatgg tgggggtcct 360  
ggccatggcg gctgcagctg ctccgcctcc cgtgaaggac tgcgagattg agccatgcaa 420  
aaagcgaaag aaagatgatg acacatctac ctgcaaaaca attacaaaat atttatcacc 480  
actagggaaag actagagaca gggttttgc tccaccaaaa cctagtaata ttctggatta 540  
ttttagaaag acttcaccca caaatgagaa gacacaattt gggaaagagt gcaagataaa 600  
gtcacctgaa tcagtacctg ttgacagcaa caaagactgt acgacaccc ttgaaatgtt 660  
ctcaaatgtt gagttttaaga agaaaagaaa gagggttaat ttatctcatc aactaaataa 720  
tattaaaact gaaaatgaag ctccaatttga aatttagtagc gacgatagca aagaagacta 780  
tagtttaat aatgatttttgg tggaaagtag tacttctgtt ttacgttaca agaaacaagt 840  
agaggtactt gcagaaaaca ttcaagatac aaaaagtcaa ccaaatacta tgacccctt 900  
gcaaaattct aaaaaagtaa atcctaaaca agggaccaca aaaaatgact tcaaaaagtt 960  
gagaaaaagg aaatgcagag atgttagtga tctatctgaa agcttaccct tggcagagga 1020  
actaaatttg cttaaaaaaag atggtaaaga tactaaacag atggagaata ctacaaggca 1080  
tgcaaaactct agagataacg taactgaagc agcccagttt aatgatagta taataactgt 1140  
ctcatatgag gaatttttaa aaagtccaa gaaaaataaa gtggaaagaga taccagactc 1200  
tacaatgtca atttgtgttc cttctgaaac tgtcgacgaa atagtcaaaaa gtggttatat 1260  
aagtgaatca gaaaactccg aaatttccca gcaggtacgc tttaagacag ttactgttct 1320  
tgcacaggtt caccctattt cggccaaaaa gacagggaaa ataccccgaa ttttcttggaa 1380  
acaaaagcaa tttgaaatgg aaaatagttt atctgatcct gagaatgaac agacagttca 1440  
gaaaagaaaaa tctaatttttg ttatcagga ggaagaatttta gaattggctg ttttggaaagc 1500  
tggaagttct gaagctgtga aacccaaatg cactctgaa gaaagacagc aattttatgaa 1560  
agcattttagg cagccagcat cagatgcact taaaaatgga gttaaaaagt cttctgataa 1620  
gcagaaagac cttaatgaaa aatgtctata tgaagtagga agagatgata attctaaaaa 1680  
aatcatggaa aattctggta tccaaatggt ttcaaaaaat ggcaatttac agttacacac 1740  
tgataaaagga agttttctga aggagaaaaaa taaaaagcta aagaagaaga ataagaaaaac 1800  
attagatact ggggcttattt caggccaaaaa cagagagggaa aacactcaaa agaaagaaaac 1860  
aacctttttc taaaaagaga aacaatatca aatagaatg agtttaagac aaaggaaaaac 1920  
agagtttttc aaaagcagca ctttattttaa caatgaaagt cttgtttatg aagatatacg 1980  
aaatgatgac cttctaaagg tttccctctct gtgtacaat aataaattgt caagaaaaac 2040  
cagcataccca gttaaaagata ttaagcttac acagtctaaa gctgaatctg aagccagctt 2100

gctaaatgtt tccacgccc agtcaactag aagatctgga agaatttagca gcacacctac	2160
tacagaaaacc attagaggta ttgattctga cgatgtacaa gataatagtc aactaaaggc	2220
ttccactcaa aaagcagcca acttatcgga aaagcacagc ttatatacag cagaattaat	2280
aacagtaccc tttgattcag agagccctat tagaatgaaa ttcaccagaa ttagtactcc	2340
caaaaaatct aagaaaaaat ctaacaaaag atctgagaaa tctgaagcaa ctgatggagg	2400
ttttacttct cagattagaa aggcaagcaa tacttcaaaa aacatatcaa aagcaaaaca	2460
attgattgaa aaagcaaaag ctttacacat cagtaggtca aaggtgactg aagaatagc	2520
gataccctta aggcgctcct ctagacatca gacactcct gaaaggaaga aattgtcaga	2580
aacagaagat tctgtataa taatagattc aagtctact gctttaaagc atccagagaa	2640
aaatcagaag aaacttcagt gtttgaatga tgtgcttagga aaaaaactta acacatccac	2700
taaaaatgtt cctggaaaaa tgaaagtgc tcctttattt cttgtcagaa aagcacaaaaa	2760
agcagctgat cctgtcccta gtttgatga aagcagtcaa gatacatctg aaaaatctca	2820
ggattgtgat gttcaatgta aagcaaagcg tgacttccta atgagtggtt tgccagattt	2880
gttgaacgg caaattgcaa agaaagctgc tgcgctggat gtgtacaatg cagttagtac	2940
cagttccag agagtcgtac atgtgcaaca aaaggatgat gggtgttggtt tgtggcattt	3000
gaaaccaccc tcttgcctc tcttaactaa atttaaagaa ctgaacacta aagtaataga	3060
tctctcaaaa tgtggatttgc ctcttggtga attttcaaca ttgaattcaa agttaaaaag	3120
cgttaactct gctgctgtgt tcatgaggac aaggaaggaa tttactgaag aagtaagaaa	3180
tctttgctt gagaaattta ggtggtcaaa tcctgaattt tcattgaaaa aatattttcc	3240
cttactccta aaaaaacaaa ttgagcacca agtacttct tccgagtgatc atagtaaaca	3300
agaactggag gctgatgtca gccataaaga aaccaaaagg aaactcgtag aagcagaaaa	3360
ttcttaagtca aaaagaaaaga aaccaaattgtt gtattcaaaa aatctggaga agaccaatag	3420
gaagtcagaa gaacttagca aaagaaacaa ctcttctggg ataaagctag attttccaa	3480
agattctgga actgaagaca tgctttggac agaaaagtat caacctcaga ctgccagtga	3540
acttata>tagga aatgagtttag ctataaaaaa gttacatagt tggttgaaag actggaaaag	3600
aagagctgaa ttggaagaaaa ggcagaatct gaagggaaaa agagatgaga aacatgaaga	3660
tttctcggtt ggcatacgtact ttaaaggcag ttcagatgtt gaagaagaga gtcgtctttg	3720
caataactgtc cttataacag gccaacagg agtggaaaaa actgctgcag tgtatgcttg	3780
tgcccaggag cttggattta agatattgtt agtgaatgcc tcttcccagc gcagtggtag	3840
acaaattcttca tctcagttga aggaagctac tcagtccttca caagtagaca aacaagggtgt	3900
aaactcacaa aaaccctgtt ttttaatag ctactacata ggcaagtcac caaaaaaaaaat	3960

aagctccct aagaaagttg ttacatcacc aagaaaagtt cctccaccat caccaaaaag	4020
tagtggacca aagcgagcac ttcctccaa aaccttgca aattatttt aagtatctcc	4080
caaacctaaa aataatgaag aaataggaat gcttctggaa aataataaag gaataaaaaa	4140
ttctttgaa cagaaacaaa ttactcagac taaatctaca aatgcaacta attcaaatgt	4200
caaagacgtt ggagctgaag aacccagcag aaaaaatgca acatctctta ttcttttga	4260
ggagggtttagt gtaattttt atgaagatgc tgggttttg aatgcaatca aaacattcat	4320
ggcaacaact aaacgacctg taatccttac tacaagtgac ccaacattta gttaatgtt	4380
tgtggctgc tttgaagaaa tcaagttcag tactccttcc ctgctaaatg ttgccagcta	4440
cctacaaatg atttgcttaa ctgagaatt tagaactgat gtaaaagact ttgtaacctt	4500
gttaactgca aatacttgt atatcagaaa aagtatcctt tacttacaat tctggattag	4560
aagtggaggt ggagtttag aagaacgacc attaaccctt taticgtggaa atagcagaaa	4620
tgtacaacta gtttgctctg aacatggcct tgataacaaa atttacccta aaaatactaa	4680
aaagaaacgt gtagaccttc caaaatgtga cagtgctgt gctgagacct tggtggcct	4740
taagaacatt tttccccat ctgaagactt atttcattt ttaaagcaca aaatcacaat	4800
gaaggaagaa tggcataaat tcatccagct tc当地tacagaa ttccaaatgc ggaatgtaga	4860
tttttatat agtaatctt agtttattct accattacca gttgatacca ttccagaaac	4920
taaaaactt tgtggccat cagtaactgtt ggatgccagt gcagcaacaa aaagtatgaa	4980
ttgtcttgct aggaaacact ctgaaagaga acagccattt gaaaagtccc agaaaaagaa	5040
acaaaagaaa acatttgtaa tattagatga tagtgcata tttgacactg acttggactt	5100
tcctgatcaa tctattagcc tgcctctgt atcatcttcc tcaaattgcag aagaaagcaa	5160
aaccggagac gaagaaagca aagccagaga caaaggaaac aatccagaga caaagaaatc	5220
tattccttgc ctcctaaaa caactgcagg aaaaaatgt tctgcccctt tttcttatttgc	5280
ttaaattct ctctctgagt tc当地ggataa catgccttc ttagatgcac ttttaactga	5340
tgtaaggaa caaaacaaaat acggtagaaa tgactttgtt gggacaaatg gaaaggttac	5400
aagtggactt tgtgtgagtt tagtcttgc gagtaatgtt ggatggactt ctcaaagctc	5460
tggagaatta aaggcagctg cagaagctct cagctttact aaatgttctt ctgttatttgc	5520
aaaagcattt gaaaccttgc attcttgc当地 gaaatttagga agagatccaa ccaacgatct	5580
tacttttat gtttcacaaa agcgcaataa tgtatacttt agtcagtcag cagcttatttgc	5640
agacaatgtt tggaagagga tatcagtc当地 taaaagtgtt tttcgagtc gatctttct	5700
ctatgtgggt aatagacaag ctatgtataat tgaataccctg ccaacccttc gaaacatctg	5760
taagactgag aagctaaaag aacaaggaaa aagtaaaaaga agattcctgc actatTTG	5820

aggaattcat cttgacattc caaaagagac tgtgaatact ttggcagctg acttccctta	5880
atgttccata ctaacaatgc tttgtataga ttatcatgtg gtccttaaga tacattttta	5940
tattatgtgg atcttcatgg aaaagtatat ttctcgatgt acatttaaa caaacaattt	6000
gtatattttt ttattggcggt gtaaatattt aaaatattt agttacaaat tttatataatg	6060
attgtatattt tttttctgaa tttttgtat tatctgattt agctttgtt gagtattttt	6120
tgtatgtgag tgaactgttt ctggaaggta gagttcatta agatgaactc cctattcaaa	6180
tggttatatat tatatatttag cttaatattc agatacatta tcttggctgc taacatttagt	6240
	6245
gtcac	

<210> 14  
 <211> 5185  
 <212> DNA  
 <213> Homo sapiens

<400> 14	
gtccttgctt ctttttcat tgatgtaaat ttttttaact taatatgaaa actctccaag	60
tcttgatcat cagccaggat tttgccacag caatttcatt ctcttacgg agcttgtatg	120
tgtcaaagta tacttcagag tgcaaacatt aagcaagtaa tatattttac ctacagatac	180
ttatgaagac attcttcgc ttcgtgagt aaaagtatat aaagctgttt ctaactgaaa	240
atgcttgaaa taagcctgta atagaaatat ttttcattt tttaaaaaaag ggcggaattc	300
ccagactgtc ttgccttct tgcacttcgc gggagaagtt gttggcgcga atggatcctg	360
agcctcgata acagattcct caaccggccc acccgccagc cagccagcgc cttcatcctg	420
gggctgcgat ggacattcgg aaattcttg gagtaatacc aagtggaaag aaacttgtaa	480
gtgaaacagt aaagaagaat gaaaaacaa agtctgtatga agaaaacttta aaagcaaaga	540
aaggaataaa ggaaatcaag gtaaatagct cccgtaaaga ggatgacttc aaacaaaagc	600
aaccaagcaa gaaaaagagg atcatctatg attcagattc agagtcagag gagacgttgc	660
agttaaaaaa tgccaaaaag ccaccagaaa aactgccagt atcttctaaa cctggtaaaa	720
tttcacggca ggatcctgtt acatacattt cagaaacaga tgaagaagat gactttatgt	780
gtaagaaggc ggcctctaaa tcaaaagaga atggaagatc tacaaatagt catttggaa	840
catcaaacat gaaaaagaat gaagaaaaca ctaagaccaa gaataaggct ttatcaccaa	900
taaaacttac acccacatca gtacttgatt attttggAAC tggaaagtgtc caaagatcta	960
ataagaagat ggtggcaagc aaaagaaaag agcttcaca aaatacagat gagtctggat	1020
taaatgtatga agccatcgcc aagcaattac agcttgatga agatgcggag ctggagaggc	1080
agttgcgtatga agatgaagag tttgccagaa cattagccat gttggatgaa gaacccaaga	1140
ccaaaaaggc tcgaaaggac acagaaggcg gagaaacgtt ttcatctgtc caagccaatt	1200

taagtaaagc agaaaaacat aaatatcctc ataaagtaaa aacagcacaa gtttcagatg	1260
aaagaaaagag ctacagtcct aggaagcaaa gtaaatatga aagttcaaaa gaatctcagc	1320
aacattccaa gtcatcagct gacaaaatag gagaagtctc ttctcccaag gccagttcta	1380
agctggcaat tatgaaaaga aaagaagaga gctcttataa agaaatagag cctgtggcct	1440
caaaaagaaa agaaaatgcc attaaattga aaggagagac aaaaactcct aagaaaacca	1500
aaagttctcc agctaaaaaa gagtcgtaa gtcctgaaga ttctgaaaag aaacgcacta	1560
attatcaagc ttatcgaagc tacttaaattc gagaaggtcc caaggctctg ggctccaaag	1620
aaataccaaa gggagctgaa aattgcttgg aaggccttat attttaatc acaggcgtgc	1680
tggagtctat tgaacgagat gaggccaagt ctctaattga acgttatggg ggaaaagtaa	1740
cagggaaatgt cagcaagaaa acaaattatc ttgtcatggg tcgtgatagt ggacagtcca	1800
agagtgataa ggccgcagcc ttggggacaa aaattattga tgaagatggc ctgttgaatc	1860
tgattcggac tatgccaggc aagaaatcca agtataat agcagttgaa actgagatga	1920
agaaaagagtc caaactggag agaacacccc aaaaaaatgt ccaaggaaaa agaaaaatta	1980
gtccatctaa aaaggaatca gaatctaaaa agagcaggcc gacttccaaa agggacagtt	2040
tggcaaagac aataaaaaag gaaacagatg tggtttggaa aagcctggat ttcaaggagc	2100
aggtggctga ggagacaagt ggtgacagca aggcttagaa tttggctgat gacagcagtg	2160
aaaacaaaagt ggaaaatttg ctctgggtgg ataaatataa gccaacctcg ctcaagacca	2220
taattggaca gcaaggtgac cagagctgtg ccaacaaact cctacgctgg ctccgaaact	2280
ggcaaaagag ttcttccgaa gataaaaaac acgcagcaaa gtttggtaaa tttccggca	2340
aagatgatgg ctctagttt aaagcagcgt tgctgtcagg ccctcctggt gttggcaaaa	2400
ccaccacagc ttccctggtg tgtcaggagt tgggatacag ctacgtggaa ctgaatgcaa	2460
gtgacaccccg gagtaagagc agtttgaagg cgattgtgc tgagtcaactg aacaatacca	2520
gcatcaaagg cttttattca aatggaggcag cctcttcagt aagcacgaaa catgctctca	2580
tcatggatga agtagatggc atggcaggca atgaggatag gggaggaatt caggaattaa	2640
ttggcctgat aaaacatact aaaattccca ttatttgtat gtgcaatgat agaaaatcatc	2700
ccaagattcg ctctctgggtt cattattgtt ttgatcttcg tttcaaaaga cctcggggtt	2760
aacagattaa gggtgctatg atgtctattg cattaaaga aggtttaaag attccccctc	2820
cagctatgaa tgaataattt tgggagcca atcaagatat cagacaggtt ttacataatc	2880
tgagtatgtg gtgtgcacga agtaaagcat taacctatga ccaggccaaa gctgattctc	2940
acagagccaa aaaggatatc aaaatgggcc catttgcgt tgcccgaaa gtgtttgcag	3000
ctggagagga gactgctcac atgtcacttg tggacaagtc agatctcttt tttcatgatt	3060

attcaatagc acccctttc gtccaggaaa attacataca cgtgaaggct gtagcagcag	3120
ggggtgacat gaaaaagcac ctgatgctt taagcagagc agcagacagc atatgcgtg	3180
gtgacctagt ggacagccag atccggagta agcaaaactg gagtcctctg cctgcgcagg	3240
ccatttatgc cagtgttctt cctggagagt tgatgagggg gtacatgacc cagttccca	3300
ccttcccaag ctggctgggg aagcactcg tctacaggcaa acatgatcgt attgttcagg	3360
acctggcctt gcatatgagt ctcagaacct actccagcaa aaggactgta aacatggatt	3420
atctgtcgct tctaaggat gcacttgtac agcccttgac ctcacaagga gtagacggag	3480
tacaggatgt tggtgcactt atggacacat attatttcat gaaagaagac tttgagaata	3540
tcatggaaat cagcagctgg ggtggcaaac ctagtccctt ttcaaagctg gatcccaagg	3600
tgaaaggcagc cttcacaaga gcttacaata aggaagccca ctttactcca tactcacttc	3660
aagctataaa ggcatctaga cacagcaca gcccattccct ggattcggaa tacaatgaag	3720
aattaaatga agatgactct caatctgatg agaaagacca agatgctata gaaactgatg	3780
ccatgatcaa gaaaaagaca aaatcttcaa agccttcaaa accagaaaaa gataaggagc	3840
ccagaaaaagg aaaaggaaaa agttcgaaga aatgaaacca tttttacta gcgacagccca	3900
cttttactc tccctcctga ccagtccagc tggcttagag aaagccttgt tttttccag	3960
agcaaccatg ttttagcata atggatgac ctgggtccc attataata aagggtggta	4020
tggctagaag ggtatgagca gtaggcttat gtacacctct tatagaggtt gataggactg	4080
cttgggtcct ccactgtcct ctgtcaatct agttagacgt gcttctgaat gactgtagaa	4140
ttggaactag aaactacacc tgggctttgg agtcagattt tagttaacaa taatgagcct	4200
ggagcagtag tactaaggcg tctttttag gcttaagaat ttatcctaatt ggccttata	4260
ggaccaatgc tgatttttt taaaggcagt tcctattatg tggtaaattt ttgtaaataa	4320
atgattatac aaaacagtca ccacctagaa ctgggtattc tttgtacatt gtcagatata	4380
tttcaaagta taaatttagga ataaaaatgt tccacatgat atgtgtacaa attttttta	4440
agattgtcat cttgtacagc aaaatattat gttggtatatt tacttccta ttaatttgca	4500
aatgatttggaa taaaaaaagc tcactgtatt ctttactaac ctgttgcgtg aaacataaca	4560
ttggcagtcg aggagtcaga actcggattc ccagcctcac tccctctgtg tgtgttgc	4620
acgtctctgc tgtgaatagg atttcatgca gtggctgtt atggggcttt gcagcgtggg	4680
gggctgcaga agattctcag acattctt ctttctggag tctcaacccaa tcaggttcag	4740
accagtagga accaggctgg gtcaggctct taatttcaact acggtggggg aaaaaatgaa	4800
cactggcttt tgttcactt gcccctgaga gatctagtc tgcctgtga tggcttccac	4860
aaaccctgtt ggaagtttgg gttattccta taactgtatg tggtgagtgg ccgtgacgta	4920

tggaccttagc ctggaattta gatgtctagc tatttttaa tccttgggt aaatgcttag	4980
aggctgtgc tacctctgtg aggtatatac taaaatgaat gtaaaaataag gatcgattc	5040
cctgcttaa ggaactggta gtcttgggt ggtgatgaga caaagttctt gagacattn	5100
aactgtattt accagtgtga aaaatgtcat ttatggagat taattcatta tgaaataaa	5160
acattgccta agcccttgcc tgctg	5185

<210> 15  
<211> 1429  
<212> DNA  
<213> *Homo sapiens*

<400> 15  
gccacgaagg ccggagagcc ggaaccggag tcgcagcggc ggagacccct gtgcggtgcg  
gagggggcgg cgccccgac tctgaccgc gcccgggtg ggccatggcg gagatcagcg  
acctggaccg gcagatcgag cagctgcgtc gctgcgagct catcaaggag agcgaagtca  
aggccctgtg cgctaaggcc agagagatct tggtagagga gagcaacgtg cagagggtgg  
actcgccagt cacagtgtgc ggcgacatcc atggacaatt ctatgaccctc aaagagctgt  
tcagagtagg tggcgacgtc cctgagacca actacctttt catggggac tttgtggacc  
gtggcttcta tagcgtcgaa acgttctcc tgctgctggc acttaagggtt cgctatcctg  
atcgcatcac actgatccgg ggcaaccatg agagtcgcca gatcacgcag gtctatggct  
tctacgatga gtgcctgcgc aagtacggct cggtgactgt gtggcgctac tgcactgaga  
tctttgacta cctcagcctg tcagccatca tcgatggcaa gatcttctgc gtgcacgggg  
gcctctcccc ctccatccag accctggatc agattcggac aatcgaccga aagcaagagg  
tgcctcatga tgggccccatg tgtgacctcc tctggtctga cccagaagac accacaggct  
ggggcgtgag ccccccggaga gccggctacc tatttggcag tgacgtggtg gcccagttca  
acgcagccaa tgacattgac atgatctgcc gtgcccacca actggtgatg gaaggttaca  
agtggactt caatgagacg gtgctcaactg tgtggtcggc acccaactac tgctaccgct  
gtggaatgt ggcagccatc ttggagctgg acgagcatct ccagaaaagat ttcatcatct  
ttgaggctgc tcccccaagag acacggggca tccccctccaa gaagccccgtg gcccactact  
tcctgtgacc ccgccccggcc cctgccccctt ccaacccttc tggccctcgc accactgtga  
ctctgcccattt ttcctcagac ggaggctggg cgtggggggg gctgtcctgg ctctgctgtc  
ccccaaagggtt gtgcttcgag ggtgaggact tctctggaga ggcctggaga cctagctcca  
tgttccttctt cctctctccc cacttgaacc atgaagtttc caataatttt tttttctttt  
tttccttctt ttttctgttt gtttttagat aaaaattttg agaaaaaaaaa tgaaaaaaatt  
1320

ctataaaaag aagaaaaatg	aaaaaaaaaa	aaaaaaaaaa	aaaaaaaaaa	aaaaaaaaaa	1380
aaaaaaaaaa	aaaaaaaaaa	aaaaaaaaaa	aaaaaaaaaa	aaaaaaaaaa	1429
<210>	16				
<211>	3865				
<212>	DNA				
<213>	Homo sapiens				
<400>	16				
ggcggattgt	ctgtcggtgc	agtagctgta	ggaaggggag	gccattttcc	60
ggagtgaggg	gcaacgggtc	ggagaaaaag	aaaaaaagaa	gggctcagcg	120
ggcccggtga	cagaggggca	cagttcggc	aggcgggtga	ggtcgctgag	180
agatgttttc	cttgcgagc	acggtgcaac	cccaggttac	agttcctctg	240
tcaatgcctt	ccatacacca	aaaaacactt	ctgtttctct	cagtggagtg	300
aaaaccagca	tcgagatgta	gttcctgagc	atgaggctcc	cagcagtgag	360
acttaaggga	ccttggatta	tctgaactaa	aaattggaca	atttgatcatg	420
atctacttcc	tggatttgt	aaaggcaaaa	acatttcttc	ccattggcat	480
tctctgcaca	atccttcttt	gaaaataaat	atggtaactt	agatataattt	540
gttcctcttg	cttgtatcga	catcattcaa	gagctttca	aagcatttgt	600
agtactggcc	agtttcata	cagtctcggg	gtttaaaac	tttggaaatca	660
gtctccagtc	tacctccgag	agattagctg	aaacacagaa	tatagcgcca	720
aggggtttct	tttgccggac	agaggatcatg	atgttgagag	tttggacaaa	780
ccaaaaatat	acctgaagct	caccaagatg	cattaaaac	tggtttgcg	840
tgaaagctca	agcactcaca	caaaaaacca	atgattccct	aaggcgaacc	900
tcttcgttct	gctgcttattc	ggcatttatg	gacttctaaa	aaaccattt	960
gcttccggac	aacaacaggg	cttgattctg	cagtagatcc	tgtccagatg	1020
cctttaaca	tgttaaaggg	gtggaggaag	ctaaacaaga	attacaggaa	1080
tcttgaaaaa	tccacaaaaa	tttactattc	ttggaggtaa	acttccaaaa	1140
tagttggacc	cccagggact	ggaaagacac	ttcttgcccc	agctgtggcg	1200
atgttccttt	ttattatgct	tctggatccg	aatttcatgta	gatgtttgt	1260
ccagccgtat	cagaatctt	tttagggaaag	caaaggcgaa	tgctccttgt	1320
ttgatgaatt	agattctgtt	ggtggaaaga	gaattgaatc	tccaatgcat	1380
ggcagaccat	aaatcaactt	cttgctgaaa	tggatggttt	taaaccat	1440
tcataatagg	agccacaaac	ttcccagagg	cattagataa	tgccttaata	1500
gttttgacat	gcaagttaca	gttccaaggc	cagatgtaaa	aggtcgaaca	1560

aatggtatct caataaaata aagtttgatc aatccgttga tccagaaaatt atagctcgag	1620
gtactgttgg cttttccgga gcagagttgg agaatcttgt gaaccaggct gcattaaaag	1680
cagctgttga tggaaaagaa atggttacca tgaaggagct ggagtttcc aaagacaaaa	1740
ttctaatggg gcctgaaaga agaagtgtgg aaattgataa caaaaacaaa accatcacag	1800
catatcatga atctggtcat gccattattt catattacac aaaagatgca atgcctatca	1860
acaaagctac aatcatgccca cgggggccaa cacttggaca tgtgtccctg ttacctgaga	1920
atgacagatg gaatgaaaact agagcccagc tgcttcaca aatggatgtt agtatggag	1980
gaagagtggc agaggagctt atatttggaa ccgaccatat tacaacaggt gcttccagtg	2040
attttataa tgccactaaa atagcaaagc ggatggttac caaatttggaa atgagtgaaa	2100
agcttggagt tatgacctac agtgatacag ggaaactaag tccagaaaacc caatctgccca	2160
tgcgacaaga aataagaatc cttctaaggg actcatatga acgagcaaaa catatcttga	2220
aaactcatgc aaaggagcat aagaatctcg cagaagcttt attgacctat gagactttgg	2280
atgccaaga gattcaaatt gttcttgagg ggaaaaagtt ggaagtgaga tgataactct	2340
cttgatatgg atgctgctg gttttattgc aagaatacaa gtgcattgc agtagtctac	2400
ttttacaacg ctttccccctc attcttgatg tggtaattt gaagggtgtg aaatgcttgc	2460
tcaatcattt gtcacatcca tccagtttgg gttattctca ttatgacacc tattgcaat	2520
tagcatcccc tggcaaatat atttgaaaa aataaagaac ttcaggatt gaaaacagct	2580
ctttgagga atgtcaatta gttattaagt tgaaagtaat taatgatttt atgtttggtt	2640
actctactag atttgataaa aattgtgcct ttagccttct atatacatca gtggaaactt	2700
aagatgcagt aattatgttc cagattgacc atgaataaaa tattttttaa tctaaatgta	2760
gagaagttgg gattaaaagc agtctcgaa acacagagcc aggaatatacg cttttggca	2820
tggtgccatg gtcacatct gtaatcccag cacttttggaa ggctgaggcg ggtggattgc	2880
ttgaggccag gagttcgaga ccagcctggc caacgtggtg aaacgctgtc tctactaaaa	2940
tacaaaaaaaaa tagggctggg cgccgggtgct cacgcctgtt atcccagcac ttttcagagg	3000
ccaaaggccggg caaatcacct gaggtcaaga gtttgagacc agcctggcca acatggtaa	3060
accccatctc tactaaacat gcaaaaattt cctggccatg gtggcagggtg cttataatcc	3120
cagctactct gggggccaaag gcaggagaat tgcttgagcc tgggagatgg aggttgcagt	3180
gagctgagat catgccactg cactccagcc tggcaacag agcaagactc tgcctcaaaa	3240
aaaaattttaa ataaatttaa atacaaaaaa aaatagccag gtgtgggtg catgcctgga	3300
atcccagcta cttgagaggc tgaggcacga gaattgcttgc aacccaggag gtggaggttg	3360
cagtgagcca agatcacagg agccactgca ctccagcctg ggtgacagag tgagactctg	3420

tctcaaaaaa aaattaaata aattattata accttcaga aatgctgtgt gcattttcat	3480
gttctttttt ttagcattac tgtcaactctc cctaatgaaa tgtacttcag agaagcagta	3540
ttttgttaaa taaatacata acctcattct gaataatgtc cctcatttg actataactg	3600
tgcttggttt caaaagcaaa attaaacaaa aatctcagtc ccctccgaag tgaactttgt	3660
gttaccctgc gtcagaaaatg ccaagttgtg tttacttttc attcagattt tgtgaatatg	3720
aacatgctgt tataggatct acagatgaat atttaactca atagaaaaat tattttagaa	3780
cacattgtat tggtattaca accagattat attcttgacg ttgacttcat taaaattatc	3840
tacaatttcc taataaaaaaa aaaaa	3865

<210> 17  
 <211> 3482  
 <212> DNA  
 <213> Homo sapiens

<400> 17	
ccctaccgcc cccaattccg ccctgcccc gcccggcg cgctagccgc cactgaggga	60
ccgaccctat aaaggccgct ccgcgagggg tgcgcagcat tcggcagagg gcgcttcgac	120
gggctggct gtgcgcctgc gcagtgtggg tcgctcccga ttccctgccc cggccggccc	180
cgccctcggtt ccgcaccctc gccccgtct cagccgcccgc tctgccccgc agcagccagc	240
cccggtccg gcagtatgtt cagctgggtc agcaaggatg cccgcccga gaaggagccg	300
gagctttcc agacggtggc tgaggggctg cggcagctgt acgcgcagaa gctgctaccc	360
ctggaggagc actaccgctt ccacgagttc cactcgcccg cgctggagga cgctgacttc	420
gacaacaagc ctatggtgc ctcgtgggg cagtacagca cgggcaagac cacccatc	480
cgacacctga tcgagcagga cttccgggg atgcgcatcg ggcccgagcc caccaccgac	540
tccttcatcg ccgtcatgca cggcccccact gagggcgtgg tgccggcaa cgcgctcg	600
gtggaccgc ggcgcctt ccgcaagctc aacgcgtttg gcaacgcttt cctcaacagg	660
ttcatgtgtg cccagctgcc caaccccgctc ctggacagca tcagcatcat cgacaccccc	720
gggatcctgt ctggagagaa gcagcggatc agcagaggct atgactttgc agccgtcctg	780
gagtggttcg cggagcgtgt ggaccgcate atcctgctct tcgacgccc caagctggac	840
atctccgatg agttctcgga agtcatcaag gctctgaaga accatgagga caagatccgc	900
gtggtgctga acaaggcaga ccagatcgag acgcagcagc tgatgcgggt gtacggggcc	960
ctcatgtggt ccctggcaa gatcatcaac acccccgagg tggtcaggggt ctacatcg	1020
tccttctggt cccacccgct cctcatcccc gacaaccgca agcttttga ggccgaggag	1080
caggacctct tcaaggacat ccagtcactg ccccgaaacg ccgcctcag gaagctcaat	1140

gacctgatca agcgggcacg gctggccaag gttcacgcct acatcatcag ctccctcaag	1200
aaagagatgc ccaatgtctt tggtaaagag agcaaaaaga aagagctggt gaacaacctg	1260
ggagagatct accagaagat tgagcgcgag caccagatct cccctggga cttcccagc	1320
ctccgcaaga tgcaggaact cctgcagacc caggactca gcaagttcca ggcgctgaag	1380
cccaagctgc tggacacggt gnatgacatg ctggccaacg acatcgccg gctgatggtg	1440
atggtgcggc aggaggagtc cctgatgcct tcccaggtgg tcaagggcgg cgcccttgac	1500
ggcaccatga acgggccgtt cgggcacggc tacggcgagg gggccggcga gggcatcgac	1560
gacgtggagt gggtgtggg caaggacaag cccacctacg acgagatctt ctacacgctg	1620
tcccctgtca acggcaagat cacgggcgcc aacgccaaga aggagatggt gaagtccaag	1680
ctccccaca ccgtgctagg gaagatctgg aagctggccg acgtggacaa ggacgggctg	1740
ctggacgacg aggagttcgc gctggccaac cacctcatca aggtcaagct ggagggccac	1800
gagctgcccgg ccgacctgcc cccgcacctg gtgcccctt ccaagcgcag acatgagtga	1860
tggcgcggg ccccgacact gccatttgca cgcccgccg ggaggcagag acggggggag	1920
gggaagcctc accatttctc aaggtccata aagactgagc ggatgtttcc tcgcctctcg	1980
aaaaggaaaa ccaccatctt tcttttaagg ctgttcctgg gcctggcggg ggaggcaggg	2040
gtgagaggat ggaattgtgt gcacaagaac tgtggctatt ttaatatata acgttagagg	2100
ctgcgttctt tgtcggccgc tcccctgtgt gccagccctg tgtgcacggc ctctgcccc	2160
cggcctttgc tgtggctgga gctggacagt gcagcgactg cgaccgtggg ggagccaggt	2220
cgccttttg gcagctgcta ggctgaggct gcatggacag gaacaccagg caccctccgt	2280
gtgcttctga gctgagggttg cttcacggga ccgtggcttc cttcctcacc tggctctgcc	2340
tccccgtgc tctcggcga agtgggttct tgtgccttcc cctccgggc ccaggctccc	2400
cgtgcgcggg ccctgccctt tcctcccgcg ccccacggc tccgacgcgc aacccgctc	2460
agcagtcaca gaagcagggc ccagccaccc tggcttttt ttggagatc agggagtag	2520
gagaatgtct tccagaaaaa tacataagct agtttctgtt ctgtaaagtg atatcttca	2580
tacttgacca aagttcccaa taacttccca accactgttc aaaagctgtg attttgtct	2640
cccttccca ccctccagcc aaggagcagc cctgcccagg gggcattagg tgtgggtacc	2700
cggggagcac ccggttcctg gaccccgatg ttgcatttcc tggctgagga aggggtgtca	2760
tcccagctcc tgccctaccc tctcactta ctggagctt gggacgcacc ctccacagtg	2820
ggaggtggtg gtgggtggcg gtggcggggc ctcacgacag cttgggtgtg gtaagaggaa	2880
gcccggtggtt ctggctaggc tctcatgtcc agacagcggg gaccagggga aaacccagcc	2940
ccttctgtaa tcccccttca tttcctactt tccttcctcc tctgttttagc aaaggagggc	3000

agctcacttg gatgtcctta caacgcccct ggccccagg ttgagcaata agaaaaccaga	3060
accttgcggc ccagtggccc gggccagttc aggccgcctc cccctcctct gcctggggcc	3120
attgagccca gcctccaggg cccgggcgcg tttgcaggcc agtggccact gtccgggctg	3180
tatggcacc aaggcaggtg gagcaccagg taccacacag ctgggcttcc caccaggctt	3240
tcccgccggg gtctcaggga gcttctcccc agcgctgctc ggagtctgca ggaactggcc	3300
ttgttctct tagcccgta ctccatacag tattaggtga ggatggatgc gggcgctgtc	3360
cttgccggga agtcactgtt gaagttgcag tggcttgttc acacctgtgg gaagagaagt	3420
gaagactttc tccttgcatt aaaaagtctg aactgtgaaa aaaaaaaaaa aaaaaaaaaa	3480
aa	3482

<210> 18  
 <211> 3517  
 <212> DNA  
 <213> Homo sapiens

<400> 18	
gacgggaccg cgagcacagg ccgctccgcg ggcgcgttcgg atcctcgcgg gaccccaccc	60
tctcccagcc tgcccagccc gctgcagccg ccagcgcgcg ccgtcgccag ctctccatct	120
gcacgtctct ccgtgaaccc cgtgagcgt gtgcagccac catgttcagc tggctgaagc	180
ggggcggggc acggggccag cagccgagg ccatccgcac ggtgacctcg gcctctaagg	240
agctgtaccg cacgaagctg ctgcccgtgg aggagcacta ccgccttggg gccttccact	300
cgcggccct ggaggacgca gacttcgacg gcaagcccat ggtgctggc gccggccagt	360
acagcacggg caagaccagc ttcatccagt acctgctgga gcaggaggtg cccggctccc	420
gcgtggggcc tgagcccacc accgactgct ttgtggccgt catgcacgg gacactgagg	480
gcaccgtgcc cggcaacgccc ctcgtcggtt accccggacaa gccctccgc aaactcaacc	540
ctttcgaaaa cacttcctc aacaggttca tgtgtgccc gctcccta at caggtcctgg	600
agagcatcag catcatcgac accccgggtt tcctgtcggt tgccaagcag agagtgagcc	660
gcggctacga cttccggcc gtgctgcgtt ggttcgcgg ggcgcgtggac ctcatcatcc	720
tgctctttga tgccacaag ctggagatct cggacgagtt ctcagaggcc atcggcgcgt	780
tgccggggcca tgaggacaag atccgcgtgg tgctcaacaa ggccgacatg gtggagacgc	840
agcagctgat gcgcgtctac ggccgcgtca tgtggccgtt gggcaaggtg gtgggcacgc	900
ccgaggtgtt gcgcgtctac atcggctcct tctggtccc gcccctccctc gtgcccgaca	960
accggcgcct cttcgagctg gaggagcagg acctttccg cgacatccag ggcctgcccc	1020
ggcacgcagc cttcgcaag ctcaacgacc tggtaagag ggcccggtg gtgcgagttc	1080
acgcttacat catcagctac ctgaagaagg agatgcctc tgtgtttggg aaggagaaca	1140

agaagaagca gctgatcctc aaactgcccg tcatcttgc gaagatttag ctggAACATC	1200
acatctcccc tggggacttt cctgattGCC agaaaatgca ggagCTGCTG atggCGCATG	1260
acttcaccaa gtttcaCTCG ctGAAGCCGA agCTGCTGG GAACtGGAC gagatGCTGA	1320
cgcacgacat CGCCAAGCTC ATGCCCTGC TGCGGCAGGA GGAGCTGGAG AGCACCGAGG	1380
tgggcgtgca ggggggcgct tttgaggGC cccacatGGG CCCGTTGTG gagCggggAC	1440
ctgacgaggc catggaggac ggCGAGGAGG GCTGGACGA CGAGGCCGAG TGGGTGGTGA	1500
ccaaggacaa gtccaaatac gacgagatct tctacaacct ggCGCCTGCC gacggcaAGC	1560
tgagcggctc caaggccaag acctggatgg tggggaccaa gctccccaac tcagtGCTGG	1620
ggcgcATCTG gaagctcAGC gatgtggacc GCGACGGCAT GCTGGATGAT gaagAGTTG	1680
cgctggccAG ccacCTCATC gaggCCAAGC tggaaggCCA CGGGCTGCC GCCAACCTGC	1740
cccgtcgccT ggtGCCACCC tccaAGCGAC GCCACAAGGG CTCCGCCGAG TGAGCCGGC	1800
ccccCTCCCA tggccCTGCT GTGGCTCCCC AGCTCCAGTC GGCTGCACGC ACACCCCTGC	1860
tccggctcac acacGCCCTG CCTGCCCTCC CTGCCAGCT GtaaggACCG GGGTCTCCC	1920
tcctcaTAC CGCCAGACAC CCCGGTGGAA GCATTAGAG GGGACCACGG GAGGGACAAG	1980
gcttctctgt CCGCCCTTA CAACCTCCAGC CTCACGTTCA CTTAGGCACA TCACACACAC	2040
actggcacAC GCAGGCATCC ATCCATCCGT CATTATTCA AATATTATT GAGCACCTAC	2100
tatgtGCCCA GCCCTGTTCT AGGCACTGGG CATTACATA gagaacaaaa tagacaaATA	2160
catctGCCCT CATGGAAGGT GACGTTCCA GGAGAGGGCA CCTACACAGT CACGCAAACA	2220
cacactaATT CCTGGCAGGG CCCCCAGCCC CTCCCTGGC TGAGCAGCCC TGTGGCTGAA	2280
atgactAGCA gataAACAGA CCCCCCTCTG CTCCGTTCC TCCGTCCAG CCAGGCAACA	2340
CCCTCAACCG GCTCCATCAC ATCCTCAGGT CTCGGACCA TGGGGGCTC AGAGGGAGA	2400
cacacctACT GCTTCCTCAG ATGGGCCCC CCGCAGCCCC TTCCCTTGCT CGGGGAAAGC	2460
CCCCAATTCT GCCCACACCC ATTATTTC CTCCTTCCTT CCTTCTTTTC TTCTTCCT	2520
TCTTCTTTT TTGTTTGC CCCCATTCT GCCCATAACCC ATTCTTTCT TTCTTCCTT	2580
CCTTCTTTT TGTTTGC CCGAGTTCTG TCCACACCCC TTCCCTTC TGTCTGTCC	2640
TTCTTTCTT TCTTTTGA TAGAATCTG CTCTGTCGCC CAGGCTGGAG TGCACTGGTG	2700
AGATCTCAGC TCACTGCAAC CTCCACCTCC TGGTTGAAG TGATTCTCGT GCCTCAGCCT	2760
CCTGAGTAGC TGGGACTGCA GGCACGCGCC ACCACGCCA GCTAATTCTT GTATTGAGT	2820
AGAGACGGGG TTTCACCATG TTGGCCAGGC TGGTCTCGAA CTCCGCATCT CAGGTGATCT	2880
GCTCGCTCG GCCTCCAAA GTGATGGGAT TACAGGCATG AGCCACCGTG CCCGGCTCA	2940
CAACCCATTTC TTAAAAAAGG ATCCCGTAGC AGGCAGAAAA GCCCCTTCCA TCCTGCTCCT	3000

ctgatactgt gcccccttgg agatatttcc gtcctccacc cacgtgtctg tggctggAAC	3060
tgcCcAGCCT gctcctggCC ccctggAAAGC ctccccACAG ctggtaatct ggacttaagg	3120
attgctggc caccgcctct ctgcctacca ccattccata tttaagtggA gcccctacgt	3180
agaaaggccc cggggcttA ttttagtctc cttttcaggG atgtcgTggg cgggggaggg	3240
ggttcttggT gctacagccc tctccccacc cctaaaggGA cgccgacgct gtttgctGCC	3300
ttcaccacat attagtgcTT gaccctggCA ggggACCCCA tggAAAAGAT ggggaAGAGC	3360
aaaatacatg gagacgacgc accctccagg atgctcgctg ggattcccAC gcccaccACT	3420
gtccccccacc ccatggctgg gaggggcctc tgaacggAAC agtgcCCCCA cagagcgaat	3480
aaagccaagg cttcttccCA aaaaaaaaaaaaaaaa aaaaaaaaaa	3517

<210> 19  
<211> 3583  
<212> DNA  
<213> Homo sapiens

<400> 19 aactttaatt gccaagattt caccctcct cctcaagccc agattatttA tcctccctcc	60
ggcctggcT gctggatgca gcagcggctg ggcttggtcc caggagcagg gagagtgcgc	120
tcccggccct cctagccgcg tgcccgggCC atggtgccgc tgagccccgc gcttgggtga	180
ggcggcggcg cggctcggag cccggcggac cggtcctacg ggacatcttC ccctgaggag	240
gagtcttccc ctggggctgc gtgccggggg cgagcggcgg ccgcgatgtt cagctggctg	300
ggtacggacg accgcccggag gaaggacccc gaggtttcc agacggtgag tgagggctc	360
aagaaaactct acaagagcaa gctgctgccc ttggaagagc attaccgctt ccacgagttc	420
cactcgcccc ccctggagga tgccgacttc gacaacaAGC ccatggttct gctggtggc	480
cagtactCCA ctgggaagac caccttcATC aggtacctgc ttgAACAGGA ctTCCCAGGC	540
atgaggatttgc ggcctgagcc caccacAGAC tccttcatttgc cggtgatgca gggagacatg	600
gaggggatca tccctggaa cgcccctggT gtggatcccA agaaaccctt cagggAAactc	660
aacgccttttgc gcaacgcctt cttgaacagg ttctgtgtgc cccagctacc taaccctgtt	720
ctggagagca tcagcgtcat cgacacacca gggatcctct ctggggagaa gcagaggatc	780
agccgggggt atgactttgc agctgtcctt gagtggttttgc cccagctacc tgaccgcattc	840
attctgctct tcgatgcccA caaaactggAC atctctgtat agttctcaga agtcatcaaa	900
gccctcaaga accacgagga caagatgcga gtggtgctga acaaagctga ccagatcgag	960
acgcagcagc tgatgcgggt gtacggggcc ctcatgtggT ctttggggaa gatcgtaac	1020
accccagagg tgatccgggt ctacatcgGC tccttctggT cccacccctt cctcatccct	1080

gacaaccgga agctcttga ggctgaggaa caggacctat tcagggacat ccagagtctg	1140
ccccgaaatg ctgccctgcg caagctcaac gacctcatca aaaggccag gctggccaag	1200
gtccacgcct acatcatcag ctctctgaag aaggagatgc cctcggttt cggaaaggac	1260
aacaagaaga aggagctggt caacaacctg gccgagatct atggccggat cgagcggag	1320
caccagatct cacctgggaa cttcccaat ctgaagagga tgcaggacca gctgcaggcc	1380
caggacttta gcaagttcca gccgctgaag agcaagctgc tggaggtgt ggacgacatg	1440
ctggcccatg acattgccc gctcatggtg cttagtgcgcc aggaggagtc acagcggccc	1500
atccagatgg tgaagggcgg agcgttcgag ggcaccctgc acggccctt tggcatggc	1560
tatggggagg gggctggaga aggtatcgat gatgctgagt gggtggtggc cagggacaag	1620
cccatgtacg acgagatctt ctacaccctg tcaccggtgg atggcaagat cacaggcgt	1680
aatgccaaga aggagatggt gcgcctcaag ctgccaaca gtgtgctggg caagatctgg	1740
aagctggccg acattgacaa ggatggcatg ctggacgacg acgagtttgc actggccaac	1800
cacctcatca aagtcaagct ggagggcac gagctgccc acgagctgcc tgcccacctc	1860
ctgccccgt ccaagaggaa agttgccag ttagtgggtg gggggacatt cagacggca	1920
gtgttagagg aggagatggg agcggtgact acacacacac acacacacac acacacacac	1980
acaaacatgc acacacacat atgcataatct tgacattgct ctgttagtga gagaggacca	2040
tgacgcccatttgcagct gatacttgg tggcacacc tccaagttct cgggattaga	2100
aggacaagag cactcccagg cccagagtc taagcctaag tctctatcgc tctccctc	2160
tcctcgccca cccccagat accagacctg aggcaattca cttgccagca cagatggcca	2220
acccacctcc agatccccca gtgctccac accccggctc tgagcaaatg gaaaagactt	2280
ttcatttagt agacaattca cttcttttc tgtgcttccc ctatctgctt tggcttcata	2340
ataagaaatc catcaagag ctaggagatc tgagggcagg cgggcagctg cagggaggag	2400
aggtgagaaa ggaagcgtct tctagagaca ttggccagg agctctgttc tttccataatc	2460
taagcctctg tcttcttcgg caaaccttc tttgaactct gccagtattt cattttaaag	2520
aatcccagag cgggagagag aagagaaaaa aattgataag agtgagaaaa ttgtcctgt	2580
gtctattgaa aaccagtcaa ggtggttta gttcatagat tttgttagat gttcttcata	2640
cctggcctat gatgtttaga tgttcataact tgactcacat ttacccagcc ctcctgcgt	2700
accaggagct gtgttaggca ctatataac attattctat gtggccctca ctgatgcccc	2760
agggaaagtat gcattagcct tcccatttg cagttgagga ggctgagtag ctcagaagg	2820
gtttaggcga cttctgaaa ctcacagaag tcacgtgatg gagagaggat tcaaagccag	2880
ggcctcagac ctcacacacac ttgtctgtgc tatgtatgtat gcaggatccc agcattgata.	2940

cccaatgaca aactatggag aacaagcaaa gatatgcaggc cccctgcagc ctcccaggac	3000
aggctggcaa gggaggaggg ccggccagca tttggtgcc catcagtctg gccatctgtc	3060
acgtcacaga agcaaaccgt gccttctggc tctgcgcccc atattccag catcatagac	3120
atccaacagc accagcagga gagtgggcta gcctgctgga tgctgttcgt gcctgtccct	3180
gctctgcctc ccacccagtt gcctgaatca tcccagctca gatgcagcca ctgtctttg	3240
tcaagtggga cctcatactacta ttctcagaag gctaacttga gaggtttggg gccttgtcc	3300
ccagagggtc cccaggact ctgcagtgtc cttggcaa at cccactgta ctcaatgccc	3360
tacattctct tctgtggtct ctcccctggc ttgcttcatg gccactgaac caatcaactt	3420
gtatgctatg ctcctactgt gatggaaaac aaaatgagta taacttattt tatatccata	3480
ttcagactat atagagaata ttctatgcat ctatgacgtg cttactactg cagtgcattt	3540
gtcatttagtc ttcatgttaa tacagtacat ttattctttg gta	3583

<210> 20  
 <211> 2941  
 <212> DNA  
 <213> Homo sapiens

<400> 20	
gagagtcgaa cccggagcag ggtcccatcc gagcgtggac tggcgccagg atgttcagct	60
ggatggggcg gcagggggc gggcgcaac gcgctggcg cgccgacgcg gtgcagacgg	120
tgacgggcgg gctgcgctcg ctctacctgc gcaaggtgct gccgctggag gaggcgtacc	180
gcttccacga gttccactcg cctgcgctgg aggacgccga cttcgagaac aagcccatga	240
tcctgctggt gggccagtac agcaccggca agaccacccat catcagatac ctactggagc	300
aggatttccc aggcatgagg attggtccgg agcccaccac agactccctc atcgccgtga	360
tgtatggaga gactgagggc agcaccggcag ggaatgctt agtcgtggac cccaaaaagc	420
cgtttagaaa gtcagtcgc tttggaaacg ctccctgaa tcgattcatg tgctcacagc	480
tccccaatca ggtcctgaag agcatcagcg tcatcgacag cccggcatac ctgtctgggg	540
agaagcagcg catcagccga ggctatgact tctgccaggt cctgcagtgg tttgccgaga	600
gggtggacag gatcatcctg ctctttgacg ctcacaagct ggacatctca gatgaattct	660
cagaggccat caaggccttc cggggccagg acgacaagat ccgtgtcgat ctgaataagg	720
ccgaccaagt ggacacgcag cagctgatgc gggtctacgg ggccctcatg tggcccttag	780
gcaagggtcat caacacgccc gaggtactgc gcgtctacat tggctccctc tggcgccagc	840
ccctgcagaa cacggacaac cgccggctct tcgaggctga ggcccaggac ctcttcagag	900
acatccagag cttcccccag aaggcagcgg tgcgcaagct caacgacccatc atcaagcag	960
cgaggctggc caaggtgcat gcctacatca tcagctacct gaagaaggag atgccaagtg	1020

tatTTggaaa ggaaaacaag aagagagac ttatcagcag gctaccggaa atctacattc	1080
agctacagcg agaataccag atttctgcag gggacttccc tgaggtcaag gctatgcagg	1140
aacagcttga gaactatgac ttcaccaaAT tccactcgct gaagcccaag ctgatcgagg	1200
cagtggacaa catgctgacG aacaagatct cGCCCTCAT gaacctcatc agccaggagg	1260
agacgagcac gcccacgcAG ctggTgcagg gcggcgcTT cGatggcacc accgaggGCC	1320
cTTcaacca gggctacGGG gaggGTGCCa aggagggcgc cgacgaggAG gagTgggtcg	1380
TggccaaAGA caagcccgtc tacgacgacG tcttctacac tctgtcgccc atcaatggca	1440
agatATcagg tgtcaacGCC aagaaggAGA tggTgacCTC caagctgccc aacagcgtcc	1500
Tgggcaagat ctggaaGctg gCcGactgcG actgcGacgg catgcttgat gaggaggAGt	1560
TcgCGctggc caagcacCTC atcaagatca agctcgacgg ctacgagctg cccagcagCC	1620
Tgccccccca CCTCGTGCcC CCCTCGACa ggaagtccCT gCCCAAGGCC gactgaggGG	1680
TgggctgcAG aacggggTgg gAACTggggg acctgggcT caggcctgCT ccaccACTGA	1740
ctCACCGAAT gacCTTggc aaggcactgc CCTCTCTGTG CCTTGGTTc CCCATCTGTA	1800
gaATggggAG ggtggacACT ggAAactAGA TGACTTCTT cacCTCCAAA ATTCCCTTA	1860
tttCTATGAA aATATTgggg GTAGGGGGT GGATTAGGAG ATTGAAGGGT TGAGAAGAAA	1920
gagAAATTGT CCAAAGAGTC CTCAGAACCT GCCTGGAGAA ATGCGCATGG GGGTGGCCT	1980
ggtAAgtccc AGGAACCACA ggaAGTgAGC TAAGGCTCAC CCAGAGCAGC TGGTCTCCA	2040
ggCTGCCTGG GCTTTTGTc CTCACTGAACA ACTCTGGAGC AGCTGTCTGT CCCTCAGAGG	2100
gcACCTGGAG GcAGCAAAGA TTATCTTATT TATTTTATTt TGTTTCTGC TATATTAGA	2160
gtggcaaaaa AATAGAGCAG AGGGTTCTG CTGTCTCCAG ACTGTCTACC AAAGAGAAGG	2220
cgACAGATGC CTCTGGTT TgGAAGGGGG ATGCACTGTG GATCTGCCAT CCATTCTGCA	2280
gtCTCCAGCA gAGCAGGcAG TCAGGCCCCA GCGTGCCTCC ATCCCAGTGC CCCGATGACC	2340
ATCTGGTCAG CCCTCCCAAC CTCCCTTCCA GGGGGCTTAC CAGCAAAGCC ATATTGGCT	2400
AGGAATTGG AATTCAACAAC CTTTATTAAAC CCCTGGCAAA ACTTCCCAGTT TGCACTGTAA	2460
ATCACATTGA AGCCTGAAGA TTAGCTTTc TTACTGGTTT TTCCAATAAG TAATAGCAAA	2520
GCCCACCTAT CATACTAAA CTACAGTAAA GAAAAGAGGT GGTCTCTGC AGTGATCGTT	2580
TAGTGTGGCC CTCAAACATT AAATATCCCC AGGTCTCCTT GGTGGGGTGGC AGGATTAAAA	2640
TTCAATCAAAC TCCCTGCCTA GTGTGTGCAG TGTCTCGGC CCTGTGGACa CAGGTGAATG	2700
AGGGACCAGC CCTGCCCTGG GCTGTTGAGA AGAGATCAGT CCACCCAGAG TTAGACATTG	2760
TTTTTGTAGA AAACAGGCAT TTATTATGTC TAGGGTTTG TGTTTTTTT TTTCCTTAC	2820
AGGATAAAAAG CCTTTATACA GAAACAAAAAT GGAACCTTc ATATAAAACC TTTCCTTAA	2880

aaaaaaaaaaa	aaaaaaaaaaa	aaaaaaaaaaa	aaaaaaaaaaa	aaaaaaaaaaa	aaaaaaaaaaa	aaaaaaaaaaa	2940
a							2941
<210>	21						
<211>	4877						
<212>	DNA						
<213>	Homo sapiens						
<400>	21						
gccccaaacc	cggaagttag	cggcggcagc	tgcgaggctc	ggagaaacag	gcgcgcggg		60
ctccgcgcc	ggccggaccc	gggcccgaga	tcatgtatgc	gccgccaccc	ccgcccaccac		120
ggagcgagaa	gcccagatag	acgccccggc	ggccccgggt	cctggagtcc	cgccgcctgc		180
tgcccgccg	aggacccac	cccgctgcc	gcccgtatgc	tgcagtgggg	cccgccatgg		240
acagggatta	cccgcagcat	gaaccccccgc	cggcggcag	cctcctgtac	agccccgcgc		300
ccctgcagag	cgccatgctg	cactgcccct	actggaacac	tttctcgctg	ccgccataacc		360
ctgccttc	cagcgacagc	cgcccgttca	tgagctccgc	ctccttc	ggcagccagc		420
cctgcccaga	caccagctat	gccccgtgg	ccaccgcctc	cagcttgcca	ccaaagacct		480
gcgactttgc	tcaggactcc	tcctattttg	aggacttctc	caacatctcc	atcttctc		540
cgtccgtgga	ctccctgtcg	gacatcgtgg	acacgcccga	cttcctgccg	gctgacagcc		600
tcaaccagg	gtccaccatc	tgggacgata	accctgcccc	ctccaccac	gataagctgt		660
tccagctca	caggccgtt	gcaggcttcg	aggacttct	gccctccac	agcacccgc		720
ttctcgtca	ctaccaggag	cagagtgtgc	agagccagcc	agaggaggag	gacgaggctg		780
aggaggagga	ggcggaggag	ctggggcaca	cagagaccta	cgccgactac	gtgccgtcca		840
agtccaagat	cgggaagcag	cacccagacc	gcgtggtgga	gaccagcaca	ctgtccagcg		900
tcccaccc	agacatcacc	tacaccctgg	ccctgccc	ggacagcggg	gcctgtctg		960
ccctgcagct	agaggccatc	acctacgcct	gccagcaaca	cgaggtcctg	ctccccagcg		1020
ggcagcgcgc	gggcttctc	atcggcgatg	gggcggcgt	ggcaaaggc	cgacgggtgg		1080
ccggagtcat	cctggagaac	cacctgcgcg	gccggaagaa	agcattgtgg	ttcagcgtct		1140
ccaaacgac	caagtacgat	gcggagcgcg	acctgcggga	catcgaagcc	acgggcac		1200
cggtgacgc	gctcagcaag	atcaagtacg	gtgacaccac	tacctcagag	ggcgtcctct		1260
tcgcccac	ctccgcctg	attggggaga	gccaggccgg	tggccagcac	cgcaactcgcc		1320
tccggcagat	cctggactgg	tgtggggagg	ccttgaggg	cgtcatcgt	ttcagcgt		1380
gtcacaaagc	caagaatgcc	ggctccacca	agatggcaa	ggccgtgcta	gacctgcaga		1440
acaagctgcc	cctggcccg	gtggtctacg	ccagcgcac	aggtgcctct	gagcctcgga		1500

acatgatcta catgagccgc ttgggtatct gggcgaggg cacacccttc cggaactttg	1560
aggagttcct gcacgccatc gagaagaggg gcgttggcgc catggagatc gtggccatgg	1620
acatgaaggt cagcggcatg tacatcgac gccagctcg cttctccggc gtcaccttcc	1680
gcatcgagga gatcccgctg gccccagcct tcgagtgcgt ctacaaccgc gcagccctgc	1740
tgtggccga ggccctgaac gtgttccagc aggccggcga ctggatccgc ctggagtcgc	1800
gcaagtccct gtggggccag ttctggtcgg cacaccagcg cttcttcaag tatctgtgca	1860
tcgcagccaa ggtgcgcgg ctggtgagc tggcccaga ggagctggcg cgagacaagt	1920
gcgtggtcat cgggctgcag tccacggcgc aggcgcgcac gcgggagggtg ctgggggaga	1980
acgatgggca cctcaactgc ttcgtctcg ccgctgaagg cgtgttcctg tcgtaattc	2040
agaagcactt tccgtccacc aagagaaaagc gggacagagg agcgggcagc aagggaaac	2100
ggcgcacctcg gggacgcggg gccaaagccc cccggctggc gtgcgagaca gcgggcgtca	2160
tccgcatcag tgacgacagc agcacggagt cggaccctgg cctggacagc gacttcaact	2220
cctccccca gtcctggtg gatgacgacg ttgtcatcgt ttagtcagtc gggctccccca	2280
gtgacgaccg gggatccctg tgcctcctgc agagagaccc gcatggcccc gggtcctgg	2340
agcgggtgga gcggctgaag caggatctgc tggacaaagt gcgcggctg ggccgggaac	2400
tgccagtcaa caccctggac gagctcatcg accagctggg cggcccccag cgggtggcgg	2460
agatgaccgg caggaaaggc cgcgtggtgt ccaggcccga cgggacggtg gccttcgagt	2520
cgcggcaga gcaggggtctg tccatcgacc acgtAACCT cagggagaag cagcgcttca	2580
tgagcggcga gaagctcgtg gccatcatct cggaggcctc cagctcggt gtctccctcc	2640
aagccgaccg ccgtgtccag aaccagcggc gccgcgtgca catgacccttgg gagtcggcgt	2700
ggagcggcga ccgcgcacatc cagcagttcg gccgcaccca ccggtccaac caggtctccg	2760
cgccagagta tgtcttcctc atctcgagc tggccgggaa gcgcgggttc gcctccatcg	2820
tggccaagcg cctggagagt ctgggggccc tgaccacgg agaccgcgc gccacggagt	2880
cccgtgacct cagcaagtac aactttgaga acaagtatgg caccgggccc ctgcactgt	2940
tcctcaccac catcctgagc cagactgaga acaaagtgcc tggcccccag ggataccctg	3000
gaggggtccc caccttcttc cgggacatga agcaggccct gctgtctgtg ggcattggtg	3060
gccgggagtc ccggaatggc tgcctggacg tggagaagga ctgtccatc accaagttcc	3120
tgaaccgcac cctggggctg gaggtgcaca agcagaatgc cctgttcag tacttctcag	3180
acacccatcgac ccacccatc gagatggaca agcgggaggg caaatacgac atggcattcc	3240
tggacccatcgac tcccggtatc gaggagatct acgaggagag ccagcagggtg ttcctggctc	3300
ccgggcaccc gcaggacggg caggtggtct tctacaagat cagcgtggac cgcggcctga	3360

agtgggagga	cgccttgcc	aagtcgctgg	cgctgacggg	cccstatgac	ggcttctacc	3420
tctcctacaa	ggtccgcgt	aacaagccc	gctgcctgct	ggcggagcag	aaccgcggcc	3480
agttcttacac	ggtgtacaag	cccaacatcg	gccggcagag	ccagctggag	gccctggaca	3540
gcctccgccc	caagttccac	cgggtcaccg	cggaggaggc	caaggagccc	tggagagtg	3600
gctacgctt	gtcgctgacg	caactgcagcc	acagcgcctg	gaaccggcac	tgccggctgg	3660
cgcaggaggg	taaggactgc	ctgcaggggc	tgcggctgct	gcaccactac	atgctgtgcg	3720
gcgcgctgct	gchgctgtgg	ggccgcatacg	ccgcccgtcat	ggccgacgtc	agcagcagca	3780
gctacctgca	gatcgtgcgg	ctgaagacca	aggacaggaa	gaagcaagt	ggcatcaaga	3840
tccccgaggg	ctgcgtgcgc	cgggtgctgc	aggagctgcg	gctgatggat	gcccacgtga	3900
agcgcaggca	ggcgcccccc	ctgggctgccc	ccgccccgccc	cgccccgcgc	ccgctggcgc	3960
tgccttgcgg	ccccggagag	gtgctggacc	tcacctacag	ccccccggcc	gaggccttcc	4020
cggcgcccc	gcacttctct	ttccccggcgc	cgctgtccct	ggacgcccggc	cccgccgtcg	4080
tgccgctggg	caccccccac	gcccaggccg	accctgcggc	cctcgccac	caggcgtcg	4140
acatcaactt	caaggaggt	ctggaggaca	tgctgcgc	gctgcacgcg	ggccgc	4200
ccgagggcgc	gctggggag	ggcgcgaaaa	cgggggcg	ggcgggcgg	gtccccgagc	4260
ggcagagcgt	gatccagttc	agcccaccc	tccccggcgc	ccaggctcct	ctctgacac	4320
ccttaggcg	aaacatgccc	caagacacag	ggaccgtttc	tcccctagga	gcagcggtgg	4380
ggagcagggc	caaggtcccc	tgaccactgc	tcagaggagc	cctaggccct	ggccgc	4440
cttcagcgc	ccgaccggg	cccccacctg	gtcagccctg	gcggggccca	ctcaggac	4500
ctggggccg	ggcgctggca	gggcctctc	tgtgcctctc	ctcctaagta	ggaaggggct	4560
ccgggtggct	gctctggac	tggcaccca	caaggctca	gtggcccaa	acccttgaaa	4620
tccgtaaac	cgggtggtcc	caagagctag	aaactcagga	aaccccagg	gctcaggccc	4680
ccgcgtctcg	ggggctccgt	ggggcagacc	cctgctaata	tatgcaattc	tccctcccc	4740
agcccttccc	tgaccctaa	gttattgccc	gctcacctct	cccaggcccc	aggccgcgg	4800
gctggcaggg	tggccctgc	ggtttctatg	tatttatagc	aagttctgat	gtacatatgt	4860
aaaggacttt	tttaaat					4877

<210> 22  
 <211> 4179  
 <212> DNA  
 <213> Homo sapiens

<400> 22						
atgggtggagc	caggcaaga	tttactgctt	gctgcttga	gtgagagtgg	aattagtccg	60
aatgacactct	ttgatattga	tggtggagat	gcaggccttg	caactccaa	gcctaccccg	120

tcagttcagc agtcagtgcc acttagtgca ttagaactag gtttggagac cgaaggcagca	180
gttcctgtta aacaagaacc agagactgta cctactccag cactattaaa tgtgaggcag	240
cctccatcta ctacaacatt tgcgtaat caaataaaatc atcttccacc cttggatct	300
acaattgtaa tgactaaaac accacctgta acaaccaaca ggcaaaccat cacttaact	360
aagtttatcc agactactgc aagcacacgc ccgtcagtct cagcaccaac agtacgaaat	420
gccatgacct ctgcacccccc aaaagaccaa gttcagctta aagatctact gaaaaataat	480
agtcttaatg aactgatgaa actaaagcca cctgctaata ttgctcagcc agtagcaaca	540
gcagctactg atgtaagcaa tggtacagta aagaaagagt cttctaataa agaaggagct	600
agaatgtgga taaacgacat gaagatgagg agttttccc caaccatgaa gttcctgtt	660
gtaaaagaag atgatgaacc agaggaagaa gatgaagaag aaatgggtca tgcagaaacc	720
tatgcagaat acatgccaat aaaattaaaa attggcctac gtcatccaga tgctgttagtg	780
gaaaccagct ctttatccag tgttactcct cctgatgttt ggtacaaaac atccattct	840
gagggaaacca ttgataatgg ctggttatca gcattgcagc ttgaggcaat tacatatgca	900
gcccagcaac atgaaacttt cctacctaattt ggagatcgtg ctggcttctt aataggtgat	960
ggtgccggtg taggaaaagg aaggacgata gcaggaatca tctatgaaaa ttatttgtt	1020
agtagaaaaac gagcattgtg gtttagtgtt tcaaattgact taaagtatga tgctgaaaga	1080
gatttaaggg atattggagc aaaaaacatt ttgggtcatt cgtaaataaa gtttaataac	1140
ggaaaaattt cttccaaaca taatgggagt gtgaaaaagg gtgttatttt tgctacttac	1200
tcttcactta ttggtaaaag ccagtctggc ggcaagtata aaacttagtt aaaacaactt	1260
ctgcattggc gcggtgatga ctgcgttggc gtgatagtgt ttgatgagtg tcataaagcc	1320
aaaaacttat gtcctgttgg ttcttcaaag ccaaccaaga caggcttagc agtttagag	1380
cttcagaaca aattgcaaaa agccagagtt gtttatgcta gtgcaactgg tgcttctgaa	1440
ccacgcaaca tggcctatat gaaccgtctt ggcataatggg gtgagggtac tccattttaga	1500
gaattcagtg attttattca agcagtagaa cggagaggag ttggtgccat ggaaatagtt	1560
gctatggata tgaagcttag aggaatgtac attgctcgac aactgagctt tactggagtg	1620
accttcaaaa ttgaggaagt tcttcttct cagagctacg ttaaaaatgta taacaaagct	1680
gtcaagctgt gggcattgc cagagagccg tttcagcaag ctgcagatct gattgtatgt	1740
gagcaacgaa tgaagaagtc catgtgggt cagttctggc ctgctcacca gaggttcttc	1800
aaatacttat gcatagcatc caaagttaaa agggttgc aactagctcg agaggaaatc	1860
aagaatggaa aatgtgtt aattggctcg cagtcacag gagaagctag aacatttagaa	1920
gctttggaaag agggcggggg agaattgaat gattttgttt caactgccaaggtgtgtt	1980

cagtcactca ttgaaaaaca tttcctgtc ccagacagga aaaaacttta tagttacta	2040
ggaatcgatt tgacagctcc aagtaacaac agttcgccaa gagatagtcc ttgtaaagaa	2100
aataaaataa agaagcggaa aggtgaagaa ataactcgag aagccaaaaa agcacgaaaa	2160
gtaggtggcc ttactggtag cagttctgac gacagtggaa gtgaatctga tgcctctgat	2220
aatgaagaaa gtgactatga gagctctaaa aacatgagtt ctggagatga tgacgatttc	2280
aacccatttt tagatgagtc taatgaggat gatgaaagtg atccctggtt aattagaaaa	2340
gaccacaaga aaaacaaaga gaaaaaaaaag aagaaaagta tagatccaga ttcttattcaa	2400
agtgccttat tagcatcagg tcttgatca aaacgaccta gttttcatc tacaccagtt	2460
atctcacctg ctcctaacag tacaccagct aacagtaaca ccaacagtaa cagtagcctt	2520
ataacaagtc aggatgctgt ggaaaggct cagcagatga agaaagacct gcttgataag	2580
ctagaaaaat tagctgaaga cctccccct aataccctgg atgaacttat cgatgaactt	2640
ggtggccctg agaacgttgc tgagatgact ggccgcaagg gacgggttgt gagcaatgat	2700
gatggaagca tatcttatga gtcaagatct gaacttgatg tgcctgtgga aatactaaac	2760
atcacagaaa aacaacgatt tatggatgga gataagaata ttgctatcat ctcagaagct	2820
gccagctcgg gtatttcatt acaagcagat aggagagcta aaaatcaaag gcgaagagtt	2880
catatgactt tagaattacc ttggagcgt gatagagcaa ttcaacagtt cggacgtact	2940
catagatcaa accaagttac tgctcctgag tatgtcttc tgatatctga actggcagga	3000
gaacaaagat ttgcatctat tggtgctaaa agacttgaga gtttgggggc acttacacat	3060
ggcgatagaa gggcaacaga atctagagat ctgagcaggt tcaactttga taataagtat	3120
ggaagaaaatg cttagaaat tgtcatgaaa tccattgtaa acttggattc tcctatggta	3180
tcaccacctc cagactatcc tggagaattt tttaaagatg ttcgacaagg actgataggc	3240
gttggcctga taaatgtaga agatcgctcg ggaattctta ctctcgataa agattataac	3300
aacataggaa aattctaaa tagaattta ggcattggagg tgcatcagca gaatgcgtta	3360
tttcagtatt ttgcggacac acttactgca gttgttcaaa atgccaaaaa aaatggaaga	3420
tatgatatgg gaatcttaga tcttggttct ggagatgaaa aagtgcggaa aagtgtatgtt	3480
aaaaagttc tgactccagg atattcaacc tctggccacg tagaattata cacaattagt	3540
gtagagaggg gaatgtcatg ggaggaagct accaagattt gggctgagct gacaggacca	3600
gacgatggct tttacttgc attgcaaata aggaacaaca agaaaactgc catcttagtt	3660
aaagaagtga atcctaaaaa gaaactttc ttatgttatac gaccaatac tggaaagcag	3720
ctcaaattag aaatttatgc tgatctaaa aagaaatata agaaggtcgt ctcagatgat	3780
gccctgatgc actggttaga tcagtataat tcatctgcag atacttgtac tcacgttat	3840

tggcgccgca attgaaaaaaaa agcaagctt gggctagttt gtgaaatagg tcttcgttgc	3900
cgtacatatt atgtattatg tggttcagtg ctgagtgtct ggacaaaagt tgagggtgtt	3960
ctagcatctg tcagtggcac aaacgtgaag atgcagatcg tgccgctaag aacggaagat	4020
gggcaacgga ttgttaggttt gatcattccg gcaaattgtg tgtctcctct tgtaaatctc	4080
ctatcaactt cagaccagtc tcaacagctt gcggtccaac agaaaacagct atggcaacag	4140
catcaccctc agagcatcac caacttgagc aacgcata	4179

<210> 23  
 <211> 4134  
 <212> DNA  
 <213> Homo sapiens

<400> 23	
cgaagatggc ggcgtcgcgt cgctctcagc atcatcacca ccatcatcaa caacagctcc	60
agcccccccc aggggcttca ggcgcgcgc cgcacactcc tccccactc agccctggcc	120
tggccccggg gaccacccca gcctctccca cggccagcgg cctggcccccc ttgcctccc	180
cgcggcacgg cctagcgctg ccggaggggg atggcagtcg ggatccgccc gacaggcccc	240
gatccccggg cccggttgc ggtaccagct gttcagtc caccagcaca atctgtaccg	300
tcgcgcgcgc tcccgtggtc ccagcggtt ctacttcatc tgccgctggg gtcgcctcca	360
accagccgg cagtggcagt aacaattcac cgctgcctc ttcttccccg acttcttcct	420
cattttcctc tccatcctcc cttggatcga gttggcgga gagccccgag gcggccggag	480
tttagcagcac agcaccactg gggctgggg cagcaggacc tgggacaggg gtcccagcag	540
tgagcggggc cttacggaa ctgctggagg cctgtcgcaa tgggacagtg tcccggtaa	600
agaggctggg ggacgcccga aacgtaaatg caaaggacat ggccggccgg aagtcttctc	660
ccctgcactt cgctgcaggt tttggaaagga aggatgttgt agaacactta ctacagatgg	720
gtgctaattgt ccacgctcgt gatgatggag gtctcatccc gcttcataat gcctgttctt	780
ttggccatgc tgaggttgtg agtctgttat tgtgccaagg agctgatcca aatgccagg	840
ataactggaa ctatacacct ctgcataag ctgttattaa agggaaagatc gatgtgtgca	900
tttgtctgct gcagcacggc gctgacccaa acattcgaa cactgatgg aaatcagccc	960
tggacctggc agatccttca gcaaaagctg tccttacagg tgaatacaag aaagacgaac	1020
tcctagaagc tgcttaggagt ggtaatgaag aaaaactaat ggctttactg actcctctaa	1080
atgtgaattt ccatgcaagt gatggcgaa agtcgactcc tttacatcta gcagcggct	1140
acaacagagt tcgaatagtt cagcttcttc ttcagcatgg tgctgatgtt catgcaaaag	1200
acaaagggtgg acttgcctt cttcataatg catgttccata tggacattat gaagtcacag	1260

aactgctact aaagcatgga gcttgtgtta atgccatgga tctctggcag tttactccac	1320
tgcacgaggc tgcttccaag aaccgtgtag aagtctgctc tttgttactt agccatggcg	1380
ctgatcctac gttagtcaac tgccatggca aaagtgtgt ggatatggct ccaactccgg	1440
agcttaggga gagattgact tatgaattta aaggtcattc tttactaca gcagccagag	1500
aagcagactt agctaaagtt aaaaaaacac tcgctctgga aatcattaat ttcaaacaac	1560
cgcagtctca tgaaacagca ctgcactgtg ctgtggcctc tctgcattccc aaacgtaaac	1620
aagtgacaga attgttactt agaaaaggag caaatgttaa tgaaaaaaaaat aaagatttca	1680
tgactccct gcatgttgcg gccgaaagag cccataatga tgtcatggaa gttctgcata	1740
agcatggcgc caagatgaat gcactggaca cccttggta gactgcttg catagagccg	1800
ccctagcagg ccacctgcag acctgcccgc tcctgctgag ttacggctct gaccctcca	1860
tcatctcctt acaaggcttc acagcagcac agatggcaa tgaaggcagtg cagcagattc	1920
tgagtgagag tacacctata cgtacttctg atgttgatta tcgactctta gaggcatcta	1980
aagctggaga cttggaaact gtgaagcaac tttgcagctc tcaaaatgtg aatttagag	2040
acttagaggg ccggcattcc acgccttac acttcgcagc aggctacaac cgctgtctg	2100
tttagtagta cctgctacac cacggtgccg atgtccatgc caaagacaag ggtggcttgg	2160
tgccttcataatgcctgt tcatatggac actatgaggt ggctgagctt ttagtaaggc	2220
atggggcttc tgtcaatgtg gcggacttat ggaaatttac ccctctccat gaagcagcag	2280
ctaaaggaaa gtatgaaatc tgcaagctcc tttaaaaca tggagcagat ccaactaaaa	2340
agaacagaga tggaaataca ctttggatt tggtaaagga aggagacaca gatattcagg	2400
acttaactgaa agggatgct gctttgtgg atgctgccaa gaagggtcgc ctggcaagag	2460
tgcagaagct ctgtacccca gagaatatca actgcagaga cacccagggc agaaattcaa	2520
ccccctctgca cctggcagca ggctataata accttggagt agctgaatat cttctagagc	2580
atggagctga tgttaatgcc caggacaagg gtggtttaat tcctcttcatt aatgcggcat	2640
cttatgggca tggtagacata gcggcttataat tgataaaata caacacgtgt gtaaatgcaa	2700
cagataagtggc ggcgttact cccctccatg aagcagccca gaaaggaagg acgcagctgt	2760
gcgccttcct cctagcgcattt ggtgcagacc ccaccatgaa gaaccaggaa ggccagacgc	2820
ctctggatct ggcaacagct gacgatatac gagcttgct gatagatgcc atgccccca	2880
aggccttacc tacctgtttt aaacctcagg ctactgttagt gagtgcctct ctgatctcac	2940
cagcatccac cccctcctgc ctctggctg ccagcagcat agacaacctc actggccctt	3000
tagcagagtt ggccgttagga ggagcctcca atgcagggga tggcgccgcg ggaacagaaaa	3060
ggaaggaagg agaagttgct ggtcttgaca tgaatatcag ccaatttcta aaaagccttg	3120

gccttgaaca ccttcgggat atctttgaaa cagaacagat tacactagat gtgtggctg	3180
atatgggtca tgaagagttg aaagaaaatag gcatcaatgc atatggcac cgccacaaat	3240
taatcaaagg agtagaaaaga ctcttaggtg gacaacaagg caccaatcct tatttgactt	3300
ttcactgtgt taatcaggga acgatttgc tggatcttgc tccagaagat aaagaatatc	3360
agtcatgttga agaagagatg caaagtacta ttcgagaaca cagagatggt ggtaatgctg	3420
.gcggcatctt caacagatac aatgtcattc gaattcaaaa agttgtcaac aagaagttga	3480
gggagcgggtt ctgccaccga cagaaggaag tgtctgagga gaatcacaac catcacaatg	3540
agcgcatgtt gtttcatggt tctccttca ttaatgccat tattcataaa gggtttgatg	3600
agcgacatgc atacatagga ggaatgtttg gggccggat ttatggct gaaaactcct	3660
caaaaaagcaa ccaatatgtt tatggaattt gaggaggaac aggctgcct acacacaagg	3720
acaggtcatg ctatatatgt cacagacaaa tgctcttctg tagagtgacc cttggaaat	3780
ccttctgca gtttagcacc atgaaaatgg cccacgcgcc tccagggcac cactcagtca	3840
ttggtagacc gagcgtcaat gggctggcat atgctgaata tgtcatctac agaggagaac	3900
aggcataccc agagtatctt atcacttacc agatcatgaa gccagaagcc cttcccaga	3960
ccgcaacagc cgcaagcag aagacctagt gaatgcctgc tggtaaggc cagatcagat	4020
ttcaacctgg gactggatta cagaggattt tttctaataa caacatcaat attctagaag	4080
tccctgacag cctagaaata agctgtttgt cttctataaa gcattgctat agtg	4134

<210> 24  
 <211> 6189  
 <212> DNA  
 <213> Homo sapiens

<400> 24	
cgcgcgcct cgctagccga aacctgcccc gccggtgccc ggccactgcg cacgcgcggg	60
acgacgtcac gtgcgtcccc gggctggac ggagctggca ggagggggct tgccagcttc	120
cgcgcgcgc tcgtttcagg acccggacgg cggattcgcg ctgcctccgc cgccgcgggg	180
cagccgggg gcagggagcc cagcgagggg cgccgcgtggg cgccgcgcgc ggactgcgc	240
ggatccggtg acagcaggga gccaagcggc ccggccctg agcgcgtctt ctccgggggg	300
cctcgccctc ctgctcgccg ggccggggct cctgctccgg ttgctggcgc tgtgctggc	360
tgtggcggcg gccaggatca tgcgggtcg cgcgtgcgc ggccggggag cggcctgcgc	420
gagcgccgcg gccaggccg tggagccggc cgcccgagag ctgttcgagg cgtgccgcaa	480
cggggacgtg gaacgagtca agaggctggt gacgcctgag aaggtgaaca gccgcgcacac	540
ggcgggcagg aaatccaccc cgctgcactt cgccgcaggt tttggcggaa aagacgtagt	600
tgaatatttg cttcagaatg gtgcaaatgt ccaagcactt gatgtatggg gccttattcc	660



tcctttacat aatgcagcat cttacggca ttagatgt gcagctctac taataaagta	2580
taatgcgt gtcaatgcca cggacaaatg ggcttcaca ctttgcacg aagcagccca	2640
aaaggacga acacagctt gtgcttgtt gctagccat ggagctgacc cgactctaa	2700
aatcaggaa ggacaaacac cttagattt agttcagca gatgatgtca gcgctttct	2760
gacagcagcc atgccccat ctgctctgcc ctcttgcac aagcctcaag tgctcaatgg	2820
tgtgagaagc ccaggagcca ctgcagatgc tctctttca ggtccatcta gcccatcaag	2880
ccttctgca gccagcagtc ttgacaactt atctggagt tttcagaac tgtcttcagt	2940
agttagttca agtggAACAG agggtgcttc cagttggag aaaaaggagg ttccaggagt	3000
agatttttagc ataactcaat tcgtaaggaa tcttgactt gagcacctaa tggatatatt	3060
ttagagagaa cagatcactt tggatgtatt agttgagatg gggcacaagg agctgaagga	3120
gattgaaatc aatgcttatg gacataggca caaactaatt aaaggagtgc agagacttat	3180
ctccggacaa caaggctta acccatattt aacttgaac acctctggta gtggacaat	3240
tcttatacat ctgtctcctg atgataaaga gttcagtct gtggaggaag agatgcaaag	3300
tacagttcga gagcacagag atggaggtca tgcaggtgga atcttcaaca gatacaat	3360
tctcaagatt cagaaggTTT gtaacaagaa actatggaa agatacactc accggagaaa	3420
agaagtttct gaagaaaacc acaaccatgc caatgaacga atgctatttc atgggtctcc	3480
ttttgtgaat gcaattatcc acaaaggctt tgatgaaagg catgcgtaca tagtgttat	3540
gtttggagct ggcatttatt ttgctaaaa ctcttccaaa agcaatcaat atgtatatgg	3600
aattggagga ggtactgggt gtccagttca caaagacaga tcttgcata tttgccacag	3660
gcagctgctc ttttgcggg taaccttggg aaagtcttc ctgcagttca gtgcataatgaa	3720
aatggcacat tctcccccag gtcatcactc agtcaactggt aggcccagtg taaatggcct	3780
agcattagct gaatatgtta tttacagagg agaacaggct tttcctgagt attaattac	3840
ttaccagatt atgaggcctg aaggtatggt cgtggataa atagttattt taagaaacta	3900
attccactga acctaaaatc atcaaagcag cagtggctc tacgtttac tccttgctg	3960
aaaaaaaaatc atcttgcaca caggcctgtg gcaaaaggat aaaaatgtga acgaagttta	4020
acattctgac ttgataaagc ttaataatg tacagtgtt tctaaatatt tcctgtttt	4080
tcagcactt aacagatgcc attccaggtt aaactgggtt gtctgtacta aattataaac	4140
agagttaact tgaaccttt atatgttatg cattgattct aacaaactgt aatgcctca	4200
acagaactaa ttttactaat acaatactgt gttcttaaa acacagcatt tacactgaat	4260
acaatttcat ttgtaaaact gtaaataaga gctttgtac tagcccagta tttatTTACA	4320
ttgctttgtta atataaatct gttttagaac tgcagcgggt tacaaaattt tttcatatgt	4380

atgttcatc tatacttcat cttacatcg catgatttag tgatcttac atttgattcc	4440
agaggctatg ttcaagtgtt agttggaaa gattgagttt tcagattaa tttgccatg	4500
ggagccctta tctgtcatta gaaatcttc tcatttaga acttatgaat atgctgaaga	4560
ttaatttgt gataccttg tatgtatgag acacattcca aagagctcta actatgatag	4620
gtcctgatta ctaaagaagc ttctttactg gcctcaattt ctatcttca tgttggaaaa	4680
tttctgcag tccttctgtg aaaatttagag caaagtgcctt ctgttttta gagaaactaa	4740
atcttgctgt tgaacaatta ttgtgttctt ttcatggaac ataagtagga tgttaacatt	4800
tccagggtgg gaagggtaat cctaaatcat ttcccaatct attctaattt ccttaaatct	4860
aaagggggaaa aaaaaaatca caaacaggac tggtagttt tttatcctaa gtatatttt	4920
tcctgttctt ttacttggtt ttatttgctg tatttatacg caatctatac atcatggta	4980
aacttaaccc agaactataa aatgttagttt ttcaagtccc cttcaggcct cctgaatggg	5040
caagtgcagt gaaacaggtg cttcctgctc ctgggttttc tctccatgtat gttatgccca	5100
attggaaata tgctgtcagt ttgtgcacca tatggtgacc acgcctgtgc tcagttggc	5160
agctatagaa ggaaatgctg tcccataaaa tgccatccct atttctaata taacactttt	5220
ttccaggaag catgcttaag catcttggta cagagacata catccattat ggcttggcaa	5280
tctctttat ttgttgactc tagctccctt caaagtcag gaaagatctt tactcactta	5340
atgaggacat tccccatcac tgtctgtacc agttcacctt tatttacgt tttattcagt	5400
ctgtaaatttta actggccctt tgcagtaact tgtacataaa gtgctagaaa atcatgttcc	5460
ttgtcctgag taagagttaa tcagagtaag tgcatttctg gagttgttcc tgtatgtaa	5520
attatgatca ttatttaaga agtcaaattcc tgatcttgc ttgcatttt tacagctctc	5580
taataattac aaatatccga aagtcatttc ttggAACACA agtggagttt gccaaatttt	5640
atatgaattt ttcaagattt ctaagcttcc aggtttata attagaagat aatgagagaa	5700
ttaatgggggt ttatatttac attatctctc aactatgttag cccatattac tcaccctatg	5760
agtgaatctg gaattgcttt tcatgtgaaa tcattgtggt ctatgagttt acaatactgc	5820
aaactgtgtt attttatcta aaccattgtt taatgagtgtt gttttccat gaatgaatatt	5880
accgtggttc atatgttagc atggcagcat ttccagatag cttttgttt gttggaaagt	5940
tggggttttt gggggggggg gaggattttt acgttgcattt gaatagccta ctttataatg	6000
atgggaatgc tttttttttt gttttggat tttttttttt gaagtgaaat ttaacttttt	6060
gtgccagtag tactattata cccatcttca gtgtcttact tgtactgtat caaattccat	6120
accctcattt aattcttaat aaaactgttc acttgtaaaa aaaaaaaaaaaa aaaaaaaaaaaa	6180
aaaaaaaaaa	6189

<210> 25  
<211> 5814  
<212> DNA  
<213> Homo sapiens

<400> 25  
gagggcgggg aagaggcgtg acggagattc ctgaggtgta gtagcctgag gttcccttat 60  
gtggccctat agctgttact gaaggaagta gcctacgtcc acgcctacaa ctgaagtctc 120  
ttgacaaaca cctcacccct gcctccggga tgaaaggggg taacctagac ctgaatggc 180  
ttgaccatct cacaactgct cgctgtacga ccgcattcgt ggcaggtaag aagattgctg 240  
tatcaactca agaaagcagt aacttcactg tctttgtatt ttgaattgca acaacaactt 300  
tgatatcaac aatgaagcaa tgatatctaa gaacaaaaga gtatttgcca acagtcatca 360  
taatatcaag tgattgtata agcagaaaca agctgtcaca gacctgtgcg tcagctaata 420  
tatggagaat gctttcttct gatactattt acttagaggc agtttaata taaatcattt 480  
caattatatc tacatcaaataaataaaaaa tgagtgaagc ccccagattc ttcgttggac 540  
cagaagatac agaaataaaat cctggaaatt atcgacattt cttccaccat gcagatgaag 600  
acgatgagga ggaagatgat tctccaccag aaaggcagat tgtggttgga atatgttcca 660  
tggcaaagaa atccaaatcc aaaccaatga aggaaattct tgaacggatc tccttattta 720  
aatatatcac agtagtagta tttgaagagg aggttatttt gaatgaacca gtggaaaact 780  
ggcctttatg tgattgtctt atttcttcc attctaaagg atttccactg gacaaagcgg 840  
ttgcctatgc aaaactcagg aatccatttga taatcaatga cttgaatatg cagtatctca 900  
tacaagatac gagagaagta tatagtatttgc ttcaagctga aggttatttt cttcctcggt 960  
atgctattttt gaaccgtgac ccaaataatc ccaaagaatg taatctgatt gaaggaaag 1020  
atcatgtaga agtaaatggg gaagttttc aaaagccatt tgttagaaaag ccagtcagtg 1080  
cagaagatca caatgtttac atttattacc caacctctgc tgggtggta agtcaaagac 1140  
tctttagaaa gattggcagt agaagtagtg tttattctcc agaaagcaat gtacgaaaaaa 1200  
caggctcata tatatatgaa gagtttatgc ccacagatgg tactgatgtt aaggttata 1260  
cagtgggtcc agattatgcc catgctgaag ctcgaaaatc tccagcactt gatggcaagg 1320  
tggAACGAGA cagtgaagga aaagaagtaa gataccctgt tattctcaat gcacgagaga 1380  
aattaattgc ttggaaagtc tgccttgctt ttaagcaaac agtttgtggc tttgatttgt 1440  
tacggggccaa tggacagtcc tatgtctgtg atgtcaatgg cttcagttt gtgaaaaattt 1500  
ccatgaagta ttatgtatgac tgtgcaaaaa tacttgaaaa tattgtatg cgagaacttg 1560  
ctccacaattt tcatattcca tggtaatac ccttagaagc tgaagatatc ccaattgtac 1620

caactacatc	tggaactatg	atgaaactta	gatgtgtcat	agctgttata	cgtcatgggg	1680
atcgaacacc	aaaacaaaaaa	atgaaaatgg	aagtgagaca	tcagaaattt	tttgatctt	1740
ttgaaaagtg	tgatggatat	aaatcaggga	aattaaaact	caaaaaacca	aaacagttac	1800
aggaagtgct	agatattgca	cgacagcttc	ttatggagct	agggcaaaat	aatgattctg	1860
aaattgaaga	aaacaagcca	aaacttgaac	aacttaagac	tgtatttagag	atgtatggtc	1920
attttctgg	aataaatcgt	aaggttcagt	tgacctatct	ccctcatggt	tgcctaaaaa	1980
catctagtga	agaggaggac	agccgaagag	aagaaccatc	tttactttt	gttctaaaat	2040
ggggaggtga	attaactcct	gcaggcaggg	tccaggctga	agaacttgga	agacgcttca	2100
ggtgttatgta	tcctggaggt	caaggagatt	atgcaggatt	tcctgggtgt	gtttactta	2160
gattacatag	cacctacaga	catgacctca	aaatatatgc	ctctgatgaa	ggacgagtcc	2220
agatgactgc	agctgcttt	gcaaaggggc	tttagcttt	ggaaggagag	cttacaccca	2280
ttcttgttca	aatggtgaaa	agtgc当地	tgaacggct	tttggatagt	gatagtgact	2340
ctctgagcag	ttgtcagcaa	cgtgtgaagg	caaggcttca	tgaaatactt	cagaaagaca	2400
gagattttac	tgctgaagat	tatgaaaagc	ttactccatc	tggaagcatt	tctttatca	2460
aatcaatgca	tttaattaaa	aaccctgtga	agacctgtga	taaagtttat	tccttaattc	2520
agagtttgac	ttctcaaatc	agacatcgaa	tggaaagatcc	taaatcatca	gatattcagc	2580
tttaccatag	tgaaacattg	gagcttatgc	tacgttagatg	gtccaagttt	gagaaagact	2640
ttaaaacaaa	gaatggaaga	tatgatatta	gtaaaatccc	tgacatataat	gactgtataa	2700
aatatgatgt	ccagcataat	gttccttga	aattagaaaa	cacaatggaa	ttatataggc	2760
tttcgaaggc	attagcagat	attgttatcc	ctcaggaata	tggtataact	aaagctgaaa	2820
aactggagat	tgccaaaggc	tactgtactc	ctctggtag	aaaaattcgc	tcagaccttc	2880
agaggacaca	agatgatgac	actgtaaata	aacttcatcc	tgtgtattct	agaggtgttc	2940
tgtctcctga	acgtcatgtt	cgtactagat	tatattttac	cagtgaaagt	catgtacatt	3000
ctttgctgtc	tattcttcgc	tatggtgct	tatgcaatga	atcaaaggat	gaacagtgg	3060
aacgagctat	ggattattta	aacggtgtca	atgagctcaa	ctacatgact	cagattgtta	3120
tcatgcttta	tgaggatcct	aataaggatc	tttcctcaga	agaacgcttt	catgttgaat	3180
tacactttag	tccgggagcc	aaaggttgc	aagaagacaa	aaatttgcca	tctggctatg	3240
gatatagacc	agcttccaga	gagaatgaag	gcaggagacc	ttttaaaatt	gataatgatg	3300
atgaaccaca	tacttctaaa	agagatgaag	ttgatcgagc	tgtgatattg	tttaaaccaa	3360
tggtatcaga	gccaaattcat	atacacagga	agtctccact	tccaagatct	aggaagacgg	3420
ctacaaatga	tgaagagagc	cccctgagtg	tgtctagccc	agagggtact	ggtacctggc	3480

tgcattatac cagtggtgtg ggtactgggc gtcgaagacg cagatcaggg	3540
aaacaaaatca	
cttcttcccc tgtctccccc aaatcattgg ctttcacatc cagtatttt ggctcatggc	3600
aacagggtgt atctgaaaat gctaattacc tgcaaacacc aagaactctt gtggAACAGA	3660
agcagaatcc tactgttaggg tctcaactgtg cgggcgtt tagcacctcg gtgctcgccc	3720
gttcttcaag cgcacctaac ctacaggatt atgctcgta tcatcgtaaa aagctgacct	3780
cttctggctg catagatgac gccacacgac gttctgctgt taaaaggttt tctatctcat	3840
ttgctcgaca cccaaaccaat ggcttgaat tgtattccat ggtgccatct atttgcctc	3900
tagaaactct tcataatgcc ctatcttaa agcaagtgg a tgaatttctt gcttccattg	3960
cttctccatc atctgacgtt ccacggaaaa ccgctgaaat ttcccttaca gctttacgtt	4020
ccagtccaat aatgagaaaa aaagtatctt taaatacgta tacacctgca aagatccctcc	4080
caacaccacc agctacccctt aagagcacta aagcaagcag caaaccagct acaagtggac	4140
cttctagtgc agttgttcct aatacctcat ctcggaaaaa gaatataact agcaaaaacag	4200
aaacgcatga acacaaaaaa aacactggaa aaaagaaaatg aaatcttagc agaagctggaa	4260
actttttata cttataaaaa tagtgtgttc ttatgtttct ctttatgcat ttatgtgttc	4320
actttaaaat gtttttaat ctaaggttt ctttgtttat gttcaggtaa ggaactgttg	4380
tcatgatctg gaaatgttta aaacaatgtt tgtagcatt ctgtgagcag caaaacttat	4440
agtgataaaaa atcgattgtt gttatatga tggttaccat gtgcacatag taatgaaaag	4500
gaacataaaaa gcccagcagg ctcgtaccaa agtcacagca gtaatgctat gtactgcaga	4560
gtctgatgag ctggcctttg tgcacacttt tattttcatg ggattgcattc ttagctgtta	4620
aaacttcttag attgaaattt gacagccagg gttacatatt ggggactttt aaagtgtctt	4680
tccaaagaga ttccattaac cgtttagatt agaatatctt tcccaattgt tacagtgaca	4740
tatatgctgc aatatccaac aactggagta ttagccacat gggttttttt ttcaatctgt	4800
gttttgaatt tttttattgt gtgttattta aaatattaca tatgcagctg ggagaactac	4860
acctttgtgc acatagattt atatattaat ttgtgaaaaa tattttcttt atatattcc	4920
ttaccataca aggtgccttg ttcatcagga aaactttgt tttgtatttt gacaagaaaag	4980
gcaccttcag agttttttt taagtatagt tgacaagtgt ataaatgtta cacttacttt	5040
cagagttctt tttagatcta aagaagtcag ttcaaaaaatg gaaatcaaca atgttaggag	5100
aaatctgaat tctgttaagt tagtaagtat tatgtatagc atctgtttt aaccatttcc	5160
attcttatcc ctatgtatc agttgatcac actaagaaaag cttaaagatt gagcatttga	5220
aataaatgct ctttataaaat gattagattt ttgaaggat attgaaatca ttgcgtgtg	5280
atttcatctg tgatgtgaaa aatcaattta ttatccttgg tgctttcccc cccaccaatg	5340

cacaataat tgtgaacagc ttgaaatgac taaaactgta atccaaatgg gacaatctga	5400
taagaatttc atgcattgg agttaaataa cttaaattgc taaagcttta gcttcaaatt	5460
tatgtttaga aaaactattg atttccatg gtatgaatat aactattgta attcttcaaa	5520
tgagactctt ctcaccctaa atagtcata attaattaac ttataggaaa taagcatact	5580
atatgttagc tgtttaaaaa ggtaccagat gtaagagtca taaatatacg caattaaaga	5640
agttcataga tttcacacga atgttaatgt gttatatacg gacatgtctt gtaaacagtt	5700
aatgttatgt aagtttctg tttgtaaaaa tgtatgttaat gtactcactg tggaggtcat	5760
aaggaagcta cttttttttt aaagtggAAC ctaataaaaa tatttcccAG aatc	5814

<210> 26  
 <211> 5544  
 <212> DNA  
 <213> Homo sapiens

<400> 26	
gccccactca agagtagcct tcctcgagga cctgccttc ccatttgctg cctgaagttt	60
atgtttcttgc ctggccaaat cagggacatg ccggcattag cgggatgagt ggggtttccg	120
gcagggatgt ggtcattgac gccagtgag ggccagagta ccacggccca ctttttcctt	180
ggagctggag atgaggggct gggcacccgt ggaataggca tgaggccaga agagagtgc	240
agcgagctcc ttgaggatga ggaggatgaa gtgcctcctg aacctcagat cattgttggc	300
atctgtgcca tgaccaagaa atccaagtcc aagccaatga ctcaaattctt agagcgactc	360
tgcagatttgc actacctgac ttttgtcatt ctgggagaag atgtatcct taatgaaccc	420
gtggaaaact ggcatttcctg ccactgcctc atctttcc actccaaagg ctttcctctg	480
gacaaagctg ttgcttactc caagcttcga aacccttcc ttatcaatga tctggccatg	540
cagtattaca tccaaatggc gagggaggtg taccggatcc tgcaggaaga gggattttgt	600
ctgcctcgat atgctgtgct caaccgtat cctgcccggc ctgaggaatg caacctgata	660
gaaggtgaag accaagttaga ggtcaatggc gctgttttc ccaagccctt tttggagaag	720
ccagtgtatc cagaagacca caatgtttac atctactacc ccagctcagc tggaggagga	780
agccagcgctc tcttcgtaa gattggcagc cgaagcagtgc ttactctcc tgagagcagc	840
gtccgaaaga cgggtcgta catctatgag gagtttatgc caacagatgg cacagatgtc	900
aagggtata cagtggggcc agattatgcc catgtgaag ctagaaaatc tccagctttg	960
gatgggaagg ttgaacgaga cagtgggggg aaagagattc gatatccagt catgctgact	1020
gccatggaaa agctggcgc cagggaaatgc tgcgtagctt tcaagcaaac agtttggaa	1080
tttgccttc ttctgtccaa tggcattcc tttgtgtgtg atgtcaatgg ctttagttt	1140
gtcaagaact cgatgaaata ctacgtgac tgcggcaaga ttctggggaa caccataatg	1200

cggagctt cccccacagtt ccagattcca tggccatcc ccacggaggc tgaggacatt	1260
ccattgttc ccaccacatc tggcaactatg atggaacttc gttgtgtcat tgcaattatt	1320
cgtcatgggg atcgtactcc caagcagaag atgaagatgg aagtgaaaca cccaaaggttt	1380
tttgctctgt ttgaaaaaca tggtggtac aagacaggaa aattaaaact caagcgacct	1440
gagcagctcc aggaggtgct ggatatcaca aggctgttgt tggctgaact ggagaaagaa	1500
ccaggtggtg agatcgagga gaagactgga aaactagagc agctgaagtc tgtactggag	1560
atgtatggtc acttctcagg tataaaccgg aaggtacaat tgacttacta ccctcatgga	1620
gtaaaagctt ctaatgaggg gcaagatcca cagagggaaa ctctggcccc atctctgttg	1680
ctggtaactga agtggggtgg agaactgact cctgctggcc gtgttcaggc tgaggagctg	1740
gggcgagctt ttgcgtgcat gtaccctgga ggacagggtg actatgctgg cttccctgg	1800
tgtggctgc ttgcgttcca tagcaacttcc cgccacgatc tcaagatcta tgcctctgat	1860
gagggtcgtg ttcagatgac tgctgctgcc ttgcccaagg gccttctggc tctagaaggg	1920
gagctgacac ccattttggt gcaaattggtg aagagtgcc acatgaatgg gctactggac	1980
agcgatgggg attccttgag cagctgccag cacccgggtga aggctcggt gcaccatatt	2040
ctacagcagg atgcaccctt tggccctgag gactacgatc agctggctcc caccagaagt	2100
acttccctgc tcaactccat gactatcatc cagaatcctg tgaaggtctg tgatcaggt	2160
tttgcctga tcgaaaacct cacccaccag atccggaaac gaatgcagga ccccaggct	2220
gtagacctgc agctctacca cagttagaca ctagagctaa tgctacagcg ttggagcaag	2280
ctggagcgtg actttcgaca gaagagtggg cgctatgata tcagtaagat ccctgacatc	2340
tatgactgtg tcaagtatga tgtgcagcac aatgggagtc tgggacttca aggcacagca	2400
gagttgctcc gtctctctaa ggcactggct gatgtggta ttccccagga gtacgggatc	2460
agtcgggagg agaaactgga aattgctgtg ggcttctgtc ttccactgtt gcggaagata	2520
ctacttgacc tgcagagaac ccacgaggat gagtctgtca acaagctgca tcccctgtac	2580
tcccaggcg tgctctcccc aggtcgccac gttcgaacgc gtctctattt caccagttag	2640
agccatgtcc actccctgtct cagtgtcttc cgttatggag gacttcttga tgagacccag	2700
gatgcacaat ggcagcgagc tttggattat ctttagtgcca tctcagagct taactacatg	2760
acccagattt tcatcatgtct ttaggaggac aacacacagg atcccttatac agaggaacgg	2820
ttccatgtgg agctacactt cagccccggg gtgaaagggtg ttgaggaaga aggcagtgcc	2880
ccggctggct gtggattccg tccagcctct tctgagaatg aggagatgaa aaccaaccaa	2940
ggcagttatgg agaacctgtg tccagggaaag gcatcagatg aaccagacccg ggcattgcag	3000
acttcacccccc agcctcctga gggccctggc cttccgagga gatcaccctt cattcgtaac	3060

cgaaaagctg gttccatgga ggtactttct gagacttcat cctcgaggcc tggggctac	3120
cggctttt catcttcacg gccacccaca gaaatgaagc agagtggcct agggttgaa	3180
gggtgttcca tgggtgcctac catctaccct ctggaaacac tgcataatgc ccttccta	3240
cgtcaagtga gtgaattctt gagtagagtc tgccagcgcc acactgatgc ccaggcacag	3300
gcatctgcag ccctcttga ttccatgcac agcagccagg cctcagataa cccattttct	3360
ccaccacgta ctcttcattc acctcccctg caactccagc agcgctctga gaagccccct	3420
tggtagaga caaggttttgc ccatgttggc caggctggtt tagagctcct gacctaagt	3480
gatctgcctg cctcggcctc ccaaagtgc gggattacag gcgtgagcca ccgcacccag	3540
ccagacagca gtggcccttc tagcaactgtg tccagtgctg gtccttcttc ccctactaca	3600
gtagatggta actcccaatt tggcttcagt gatcaaccct ccctaaattc acacgtggct	3660
gaagaacatc aaggccttgg gctgctccag gagaccctg ggagtggagc acaagagctc	3720
tccatagaag gggagcaaga gctttttgaa ccaaattcagt ccccacaggt gccacctatg	3780
gaaaccagcc agccatacga ggaggtcagc cagccatgtc aggaggtccc tgacatcagc	3840
cagccatgcc aggacatttc tgaggcgctc agccagccat gtcagaaggt ccctgacatc	3900
agccagcaat gccaggagaa ccatgacaat ggtaaccaca catgccagga gttccctcac	3960
atcagccagc catgccagaa gtccagccaa ctgtgccaga aagtctctga ggaagtttgc	4020
cagctatgtc tggagaactc cgaggaggc agccagccat gccaggggt ctctgtggag	4080
gttggcaagc tggccataa gttccatgta ggggtggta gcttggtcca ggaaaccctt	4140
gtagaagttg gcagcccagc tgaagagatc cctgaggagg tcattccagcc ataccaggag	4200
ttctctgtgg aggttggcag gctggccag gagacttctg cgatcaatct gttatctcag	4260
ggcatccctg agattgataa accatcccag gagttccctg aggagattga tctgcaggcc	4320
caggaggcctc ctgaggagat aaattagaag tcctgggtgg tccctgaagt gattgatcag	4380
ctgcctggag aggttattcc tcaagcccag catccatctg gtatccaaa ccctcagagc	4440
cagtctctag cccatgacca gcactcaccc cttccaccag caacatgtga ttaattttct	4500
cattagtggt atcacactat accagccatt tgagccagca acctttctg ttggcttaact	4560
cactggccag ctctcaccag cggtgtctgg ggaagtagtt ctctttgtat gaagcatacc	4620
tgtgccagag ctgtgggtga ggaggagcca gtttttaggtt cgaagaagcc attggctcct	4680
cacttagcca ttagacttga ataggatttc cttgggggtgg gttggctgt cattacccag	4740
ccttctctga gcatcctagg aaatcacaga ttgttaaagg aaatgccgt tcactgctga	4800
agacaccatc tggcgacagc aaatgcaaaa gaggggactc tagggtcttc actttctgg	4860
ggaaatgttc acgacttctc aaggtacgct tagaccatat gtgcattcag gggactcttg	4920

tctttgccag cactcctcag	gtgagcccg gatggctatc	tgaaaatgtct ggaaaataga	4980
gtcccctccc caaacatctg	acatggccac taaatctta	gacttgcctt atagatcctt	5040
ccatatcatc cccaagtgcc	ccttcgtcct cagtcgtt	cctctggca atagctctag	5100
ggaaaaagtag gtattagact	gctgtcaaa attcagagca	actactaaa atgctctaga	5160
ggattcttag ccagtctgtg	aaagtggcc tcctctgtgg	tggggctgtt tggcttagga	5220
gatgctgttag tagtagagat	aatcaggttc tatttaagc	aatcagcaga gatcaatatt	5280
gtacagatac aagcaaaggt	ttaataaata tatatatata	tatatttttc tgtagttgtg	5340
ccagggacag actggccaaa	gaccaaacac tgcagggtcc	ccaagaaatt tgtccttata	5400
tcattgtgtc tgagggccag	atatgattat gaagctttt	ccaaagatcc agggatggga	5460
atgggagtgg gtaggagggg	taatgcggtc attggagtcg	ggggctggaa cattatgagt	5520
gctcaataaa tataaactaa	tgag		5544

<210> 27  
 <211> 397  
 <212> PRT  
 <213> Homo sapiens

<400> 27

Met Arg Cys Pro Lys Cys Leu Leu Cys Leu Ser Ala Leu Leu Thr Leu  
 1 5 10 15

Leu Gly Leu Lys Val Tyr Ile Glu Trp Thr Ser Glu Ser Arg Leu Ser  
 20 25 30

Lys Ala Tyr Pro Ser Pro Arg Gly Thr Pro Pro Ser Pro Thr Pro Ala  
 35 40 45

Asn Pro Glu Pro Thr Leu Pro Ala Asn Leu Ser Thr Arg Leu Gly Gln  
 50 55 60

Thr Ile Pro Leu Pro Phe Ala Tyr Trp Asn Gln Gln Trp Arg Leu  
 65 70 75 80

Gly Ser Leu Pro Ser Gly Asp Ser Thr Glu Thr Gly Gly Cys Gln Ala  
 85 90 95

Trp Gly Ala Ala Ala Ala Thr Glu Ile Pro Asp Phe Ala Ser Tyr Pro  
 100 105 110

Lys Asp Leu Arg Arg Phe Leu Leu Ser Ala Ala Cys Arg Ser Phe Pro  
 115 120 125

Gln Trp Leu Pro Gly Gly Ser Gln Val Ser Ser Cys Ser Asp  
130 135 140

Thr Asp Val Pro Tyr Leu Leu Leu Ala Val Lys Ser Glu Pro Gly Arg  
145 150 155 160

Phe Ala Glu Arg Gln Ala Val Arg Glu Thr Trp Gly Ser Pro Ala Pro  
165 170 175

Gly Ile Arg Leu Leu Phe Leu Leu Gly Ser Pro Val Gly Glu Ala Gly  
180 185 190

Pro Asp Leu Asp Ser Leu Val Ala Trp Glu Ser Arg Arg Tyr Ser Asp  
195 200 205

Leu Leu Leu Trp Asp Phe Leu Asp Val Pro Phe Asn Gln Thr Leu Lys  
210 215 220

Asp Leu Leu Leu Leu Ala Trp Leu Gly Arg His Cys Pro Thr Val Ser  
225 230 235 240

Phe Val Leu Arg Ala Gln Asp Asp Ala Phe Val His Thr Pro Ala Leu  
245 250 255

Leu Ala His Leu Arg Ala Leu Pro Pro Ala Ser Ala Arg Ser Leu Tyr  
260 265 270

Leu Gly Glu Val Phe Thr Gln Ala Met Pro Leu Arg Lys Pro Gly Gly  
275 280 285

Pro Phe Tyr Val Pro Glu Ser Phe Phe Glu Gly Gly Tyr Pro Ala Tyr  
290 295 300

Ala Ser Gly Gly Tyr Val Ile Ala Gly Arg Leu Ala Pro Trp Leu  
305 310 315 320

Leu Arg Ala Ala Ala Arg Val Ala Pro Phe Pro Phe Glu Asp Val Tyr  
325 330 335

Thr Gly Leu Cys Ile Arg Ala Leu Gly Leu Val Pro Gln Ala His Pro  
340 345 350

Gly Phe Leu Thr Ala Trp Pro Ala Asp Arg Thr Ala Asp His Cys Ala  
355 360 365

Phe Arg Asn Leu Leu Leu Val Arg Pro Leu Gly Pro Gln Ala Ser Ile

370

375

380

Arg Leu Trp Lys Gln Leu Gln Asp Pro Arg Leu Gln Cys  
385 390 395

<210> 28  
<211> 377  
<212> PRT  
<213> Homo sapiens

<400> 28

Met Arg Ser Ala Thr Ala Arg Pro Arg Arg Arg Ala Arg Arg Glu Gly  
1 5 10 15

Glu Gly Gly Arg His Arg Gly Pro Pro Pro Asp Pro Ala Arg Ser Ser  
20 25 30

Tyr Pro Thr Arg Val Gln Pro Arg Arg Pro Thr Lys Gly Thr His Arg  
35 40 45

Arg Arg Pro Arg Leu Arg Asp Pro Phe Asp Phe Ala Arg Tyr Leu Arg  
50 55 60

Ala Lys Asp Gln Arg Arg Phe Pro Leu Leu Ile Asn Gln Pro His Lys  
65 70 75 80

Cys Arg Gly Asp Gly Ala Pro Gly Gly Arg Pro Asp Leu Leu Ile Ala  
85 90 95

Val Lys Ser Val Ala Glu Asp Phe Glu Arg Arg Gln Ala Val Arg Gln  
           100                 105                 110

Thr Trp Gly Ala Glu Gly Arg Val Gln Gly Ala Leu Val Arg Arg Val  
115 120 125

Phe Leu Leu Gly Val Pro Arg Gly Ala Gly Ser Gly Gly Ala Asp Glu  
 130 135 140

Val Gly Glu Gly Ala Arg Thr His Trp Arg Ala Leu Leu Arg Ala Glu  
145 . 150 . 155 . 160

Ser Leu Ala Tyr Ala Asp Ile Leu Leu Trp Ala Phe Asp Asp Thr Phe  
                   165                 170                 175

Phe Asn Leu Thr Leu Lys Glu Ile His Phe Leu Ala Trp Ala Ser Ala  
 180 185 190

Phe Cys Pro Asp Val Arg Phe Val Phe Lys Gly Asp Ala Asp Val Phe  
195 200 205

Val Asn Val Gly Asn Leu Leu Glu Phe Leu Ala Pro Arg Asp Pro Ala  
210 215 220

Gln Asp Leu Leu Ala Gly Asp Val Ile Val His Ala Arg Pro Ile Arg  
225 230 235 240

Thr Arg Ala Ser Lys Tyr Tyr Ile Pro Glu Ala Val Tyr Gly Leu Pro  
245 250 255

Ala Tyr Pro Ala Tyr Ala Gly Gly Gly Phe Val Leu Ser Gly Ala  
260 265 270

Thr Leu His Arg Leu Ala Gly Ala Cys Ala Gln Val Glu Leu Phe Pro  
275 280 285

Ile Asp Asp Val Phe Leu Gly Met Cys Leu Gln Arg Leu Arg Leu Thr  
290 295 300

Pro Glu Pro His Pro Ala Phe Arg Thr Phe Gly Ile Pro Gln Pro Ser  
305 310 315 320

Ala Ala Pro His Leu Ser Thr Phe Asp Pro Cys Phe Tyr Arg Glu Leu  
325 330 335

Val Val Val His Gly Leu Ser Ala Ala Asp Ile Trp Leu Met Trp Arg  
340 345 350

Leu Leu His Gly Pro His Gly Pro Ala Cys Ala His Pro Gln Pro Val  
355 360 365

Ala Ala Gly Pro Phe Gln Trp Asp Ser  
370 375

<210> 29  
<211> 378  
<212> PRT  
<213> Homo sapiens

<400> 29

Met Leu Pro Pro Gln Pro Ser Ala Ala His Gln Gly Arg Gly Arg  
1 5 10 15

Ser Gly Leu Leu Pro Lys Gly Pro Ala Met Leu Cys Arg Leu Cys Trp  
20 25 30

Leu Val Ser Tyr Ser Leu Ala Val Leu Leu Gly Cys Leu Leu Phe  
35 40 45

Leu Arg Lys Ala Ala Lys Pro Ala Gly Asp Pro Thr Ala His Gln Pro  
50 55 60

Phe Trp Ala Pro Pro Thr Pro Arg His Ser Arg Cys Pro Pro Asn His  
65 70 75 80

Thr Val Ser Ser Ala Ser Leu Ser Leu Pro Ser Arg His Arg Leu Phe  
85 90 95

Leu Thr Tyr Arg His Cys Arg Asn Phe Ser Ile Leu Leu Glu Pro Ser  
100 105 110

Gly Cys Ser Lys Asp Thr Phe Leu Leu Leu Ala Ile Lys Ser Gln Pro  
115 120 125

Gly His Val Glu Arg Arg Ala Ala Ile Arg Ser Thr Trp Gly Arg Val  
130 135 140

Gly Gly Trp Ala Arg Gly Arg Gln Leu Lys Leu Val Phe Leu Leu Gly  
145 150 155 160

Val Ala Gly Ser Ala Pro Pro Ala Gln Leu Leu Ala Tyr Glu Ser Arg  
165 170 175

Glu Phe Asp Asp Ile Leu Gln Trp Asp Phe Thr Glu Asp Phe Phe Asn  
180 185 190

Leu Thr Leu Lys Glu Leu His Leu Gln Arg Trp Val Val Ala Ala Cys  
195 200 205

Pro Gln Ala His Phe Met Leu Lys Gly Asp Asp Asp Val Phe Val His  
210 215 220

Val Pro Asn Val Leu Glu Phe Leu Asp Gly Trp Asp Pro Ala Gln Asp  
225 230 235 240

Leu Leu Val Gly Asp Val Ile Arg Gln Ala Leu Pro Asn Arg Asn Thr  
245 250 255

Lys Val Lys Tyr Phe Ile Pro Pro Ser Met Tyr Arg Ala Thr His Tyr  
260 265 270

Pro Pro Tyr Ala Gly Gly Gly Tyr Val Met Ser Arg Ala Thr Val

275

280

285

Arg Arg Leu Gln Ala Ile Met Glu Asp Ala Glu Leu Phe Pro Ile Asp  
 290 295 300

Asp Val Phe Val Gly Met Cys Leu Arg Arg Leu Gly Leu Ser Pro Met  
 305 310 315 320

His His Ala Gly Phe Lys Thr Phe Gly Ile Arg Arg Pro Leu Asp Pro  
 325 330 335

Leu Asp Pro Cys Leu Tyr Arg Gly Leu Leu Leu Val His Arg Leu Ser  
 340 345 350

Pro Leu Glu Met Trp Thr Met Trp Ala Leu Val Thr Asp Glu Gly Leu  
 355 360 365

Lys Cys Ala Ala Gly Pro Ile Pro Gln Arg  
 370 375

<210> 30  
<211> 372  
<212> PRT  
<213> Homo sapiens

<400> 30

Met Lys Tyr Leu Arg His Arg Arg Pro Asn Ala Thr Leu Ile Leu Ala  
 1 5 10 15

Ile Gly Ala Phe Thr Leu Leu Leu Phe Ser Leu Leu Val Ser Pro Pro  
 20 25 30

Thr Cys Lys Val Gln Glu Gln Pro Pro Ala Ile Pro Glu Ala Leu Ala  
 35 40 45

Trp Pro Thr Pro Pro Thr Arg Pro Ala Pro Ala Pro Cys His Ala Asn  
 50 55 60

Thr Ser Met Val Thr His Pro Asp Phe Ala Thr Gln Pro Gln His Val  
 65 70 75 80

Gln Asn Phe Leu Leu Tyr Arg His Cys Arg His Phe Pro Leu Leu Gln  
 85 90 95

Asp Val Pro Pro Ser Lys Cys Ala Gln Pro Val Phe Leu Leu Val  
 100 105 110

Ile Lys Ser Ser Pro Ser Asn Tyr Val Arg Arg Glu Leu Leu Arg Arg  
115 120 125

Thr Trp Gly Arg Glu Arg Lys Val Arg Gly Leu Gln Leu Arg Leu Leu  
130 135 140

Phe Leu Val Gly Thr Ala Ser Asn Pro His Glu Ala Arg Lys Val Asn  
145 150 155 160

Arg Leu Leu Glu Leu Glu Ala Gln Thr His Gly Asp Ile Leu Gln Trp  
165 170 175

Asp Phe His Asp Ser Phe Phe Asn Leu Thr Leu Lys Gln Val Leu Phe  
180 185 190

Leu Gln Trp Gln Glu Thr Arg Cys Ala Asn Ala Ser Phe Val Leu Asn  
195 200 205

Gly Asp Asp Asp Val Phe Ala His Thr Asp Asn Met Val Phe Tyr Leu  
210 215 220

Gln Asp His Asp Pro Gly Arg His Leu Phe Val Gly Gln Leu Ile Gln  
225 230 235 240

Asn Val Gly Pro Ile Arg Ala Phe Trp Ser Lys Tyr Tyr Val Pro Glu  
245 250 255

Val Val Thr Gln Asn Glu Arg Tyr Pro Pro Tyr Cys Gly Gly Gly  
260 265 270

Phe Leu Leu Ser Arg Phe Thr Ala Ala Ala Leu Arg Arg Ala Ala His  
275 280 285

Val Leu Asp Ile Phe Pro Ile Asp Asp Val Phe Leu Gly Met Cys Leu  
290 295 300

Glu Leu Glu Gly Leu Lys Pro Ala Ser His Ser Gly Ile Arg Thr Ser  
305 310 315 320

Gly Val Arg Ala Pro Ser Gln His Leu Ser Ser Phe Asp Pro Cys Phe  
325 330 335

Tyr Arg Asp Leu Leu Leu Val His Arg Phe Leu Pro Tyr Glu Met Leu  
340 345 350

Leu Met Trp Asp Ala Leu Asn Gln Pro Asn Leu Thr Cys Gly Asn Gln  
355 360 365

Thr Gln Ile Tyr  
370

<210> 31  
<211> 401  
<212> PRT  
<213> Homo sapiens

<400> 31

Met Ser Leu Trp Lys Lys Thr Val Tyr Arg Ser Leu Cys Leu Ala Leu  
1 5 10 15

Ala Leu Leu Val Ala Val Thr Val Phe Gln Arg Ser Leu Thr Pro Gly  
20 25 30

Gln Phe Leu Gln Glu Pro Pro Pro Pro Thr Leu Glu Pro Gln Lys Ala  
35 40 45

Gln Lys Pro Asn Gly Gln Leu Val Asn Pro Asn Asn Phe Trp Lys Asn  
50 55 60

Pro Lys Asp Val Ala Ala Pro Thr Pro Met Ala Ser Gln Gly Pro Gln  
65 70 75 80

Ala Trp Asp Val Thr Thr Asn Cys Ser Ala Asn Ile Asn Leu Thr  
85 90 95

His Gln Pro Trp Phe Gln Val Leu Glu Pro Gln Phe Arg Gln Phe Leu  
100 105 110

Phe Tyr Arg His Cys Arg Tyr Phe Pro Met Leu Leu Asn His Pro Glu  
115 120 125

Lys Cys Arg Gly Asp Val Tyr Leu Leu Val Val Val Lys Ser Val Ile  
130 135 140

Thr Gln His Asp Arg Arg Glu Ala Ile Arg Gln Thr Trp Gly Arg Glu  
145 150 155 160

Arg Gln Ser Ala Gly Gly Arg Gly Ala Val Arg Thr Leu Phe Leu  
165 170 175

Leu Gly Thr Ala Ser Lys Gln Glu Glu Arg Thr His Tyr Gln Gln Leu  
180 185 190

Leu Ala Tyr Glu Asp Arg Leu Tyr Gly Asp Ile Leu Gln Trp Gly Phe

195

200

205

Leu Asp Thr Phe Phe Asn Leu Thr Leu Lys Glu Ile His Phe Leu Lys  
 210 215 220

Trp Leu Asp Ile Tyr Cys Pro His Val Pro Phe Ile Phe Lys Gly Asp  
 225 230 235 240

Asp Asp Val Phe Val Asn Pro Thr Asn Leu Leu Glu Phe Leu Ala Asp  
 245 250 255

Arg Gln Pro Gln Glu Asn Leu Phe Val Gly Asp Val Leu Gln His Ala  
 260 265 270

Arg Pro Ile Arg Arg Lys Asp Asn Lys Tyr Tyr Ile Pro Gly Ala Leu  
 275 280 285

Tyr Gly Lys Ala Ser Tyr Pro Pro Tyr Ala Gly Gly Gly Phe Leu  
 290 295 300

Met Ala Gly Ser Leu Ala Arg Arg Leu His His Ala Cys Asp Thr Leu  
 305 310 315 320

Glu Leu Tyr Pro Ile Asp Asp Val Phe Leu Gly Met Cys Leu Glu Val  
 325 330 335

Leu Gly Val Gln Pro Thr Ala His Glu Gly Phe Lys Thr Phe Gly Ile  
 340 345 350

Ser Arg Asn Arg Asn Ser Arg Met Asn Lys Glu Pro Cys Phe Phe Arg  
 355 360 365

Ala Met Leu Val Val His Lys Leu Leu Pro Pro Glu Leu Leu Ala Met  
 370 375 380

Trp Gly Leu Val His Ser Asn Leu Thr Cys Ser Arg Lys Leu Gln Val  
 385 390 395 400

Leu

<210> 32  
 <211> 397  
 <212> PRT  
 <213> Homo sapiens

<400> 32

Met Ser Val Gly Arg Arg Arg Ile Lys Leu Leu Gly Ile Leu Met Met  
1 5 10 15

Ala Asn Val Phe Ile Tyr Phe Ile Met Glu Val Ser Lys Ser Ser Ser  
20 25 30

Gln Glu Lys Asn Gly Lys Gly Glu Val Ile Ile Pro Lys Glu Lys Phe  
35 40 45

Trp Lys Ile Ser Thr Pro Pro Glu Ala Tyr Trp Asn Arg Glu Gln Glu  
50 55 60

Lys Leu Asn Arg Gln Tyr Asn Pro Ile Leu Ser Met Leu Thr Asn Gln  
65 70 75 80

Thr Gly Glu Ala Gly Arg Leu Ser Asn Ile Ser His Leu Asn Tyr Cys  
85 90 95

Glu Pro Asp Leu Arg Val Thr Ser Val Val Thr Gly Phe Asn Asn Leu  
100 105 110

Pro Asp Arg Phe Lys Asp Phe Leu Leu Tyr Leu Arg Cys Arg Asn Tyr  
115 120 125

Ser Leu Leu Ile Asp Gln Pro Asp Lys Cys Ala Lys Lys Pro Phe Leu  
130 135 140

Leu Leu Ala Ile Lys Ser Leu Thr Pro His Phe Ala Arg Arg Gln Ala  
145 150 155 160

Ile Arg Glu Ser Trp Gly Gln Glu Ser Asn Ala Gly Asn Gln Thr Val  
165 170 175

Val Arg Val Phe Leu Leu Gly Gln Thr Pro Pro Glu Asp Asn His Pro  
180 185 190

Asp Leu Ser Asp Met Leu Lys Phe Glu Ser Glu Lys His Gln Asp Ile  
195 200 205

Leu Met Trp Asn Tyr Arg Asp Thr Phe Phe Asn Leu Ser Leu Lys Glu  
210 215 220

Val Leu Phe Leu Arg Trp Val Ser Thr Ser Cys Pro Asp Thr Glu Phe  
225 230 235 240

Val Phe Lys Gly Asp Asp Asp Val Phe Val Asn Thr His His Ile Leu  
245 250 255

Asn Tyr Leu Asn Ser Leu Ser Lys Thr Lys Ala Lys Asp Leu Phe Ile  
260 265 270

Gly Asp Val Ile His Asn Ala Gly Pro His Arg Asp Lys Lys Leu Lys  
275 280 285

Tyr Tyr Ile Pro Glu Val Val Tyr Ser Gly Leu Tyr Pro Pro Tyr Ala  
290 295 300

Gly Gly Gly Phe Leu Tyr Ser Gly His Leu Ala Leu Arg Leu Tyr  
305 310 315 320

His Ile Thr Asp Gln Val His Leu Tyr Pro Ile Asp Asp Val Tyr Thr  
325 330 335

Gly Met Cys Leu Gln Lys Leu Gly Leu Val Pro Glu Lys His Lys Gly  
340 345 350

Phe Arg Thr Phe Asp Ile Glu Glu Lys Asn Lys Asn Ile Cys Ser  
355 360 365

Tyr Val Asp Leu Met Leu Val His Ser Arg Lys Pro Gln Glu Met Ile  
370 375 380

Asp Ile Trp Ser Gln Leu Gln Ser Ala His Leu Lys Cys  
385 390 395

<210> 33  
<211> 384  
<212> PRT  
<213> Homo sapiens

<400> 33

Met Ala Phe Pro Cys Arg Arg Ser Leu Thr Ala Lys Thr Leu Ala Cys  
1 5 10 15

Leu Leu Val Gly Val Ser Phe Leu Ala Leu Gln Gln Trp Phe Leu Gln  
20 25 30

Ala Pro Arg Ser Pro Arg Glu Glu Arg Ser Pro Gln Glu Glu Thr Pro  
35 40 45

Glu Gly Pro Thr Asp Ala Pro Ala Ala Asp Glu Pro Pro Ser Glu Leu  
50 55 60

Val Pro Gly Pro Pro Cys Val Ala Asn Ala Ser Ala Asn Ala Thr Ala

65

70

75

80

Asp Phe Glu Gln Leu Pro Ala Arg Ile Gln Asp Phe Leu Arg Tyr Arg  
 85 90 95

His Cys Arg His Phe Pro Leu Leu Trp Asp Ala Pro Ala Lys Cys Ala  
 100 105 110

Gly Gly Arg Gly Val Phe Leu Leu Ala Val Lys Ser Ala Pro Glu  
 115 120 125

His Tyr Glu Arg Arg Glu Leu Ile Arg Arg Thr Trp Gly Gln Glu Arg  
 130 135 140

Ser Tyr Gly Gly Arg Pro Val Arg Arg Leu Phe Leu Leu Gly Thr Pro  
 145 150 155 160

Gly Pro Glu Asp Glu Ala Arg Ala Glu Arg Leu Ala Glu Leu Val Ala  
 165 170 175

Leu Glu Ala Arg Glu His Gly Asp Val Leu Gln Trp Ala Phe Ala Asp  
 180 185 190

Thr Phe Leu Asn Leu Thr Leu Lys His Leu His Leu Leu Asp Trp Leu  
 195 200 205

Ala Ala Arg Cys Pro His Ala Arg Phe Leu Leu Ser Gly Asp Asp Asp  
 210 215 220

Val Phe Val His Thr Ala Asn Val Val Arg Phe Leu Gln Ala Gln Pro  
 225 230 235 240

Pro Gly Arg His Leu Phe Ser Gly Gln Leu Met Glu Gly Ser Val Pro  
 245 250 255

Ile Arg Asp Ser Trp Ser Lys Tyr Phe Val Pro Pro Gln Leu Phe Pro  
 260 265 270

Gly Ser Ala Tyr Pro Val Tyr Cys Ser Gly Gly Phe Leu Leu Ser  
 275 280 285

Gly Pro Thr Ala Arg Ala Leu Arg Ala Ala Arg His Thr Pro Leu  
 290 295 300

Phe Pro Ile Asp Asp Ala Tyr Met Gly Met Cys Leu Glu Arg Ala Gly  
 305 310 315 320

Leu Ala Pro Ser Gly His Glu Gly Ile Arg Pro Phe Gly Val Gln Leu  
325 330 335

Pro Gly Ala Gln Gln Ser Ser Phe Asp Pro Cys Met Tyr Arg Glu Leu  
340 345 350

Leu Leu Val His Arg Phe Ala Pro Tyr Glu Met Leu Leu Met Trp Lys  
355 360 365

Ala Leu His Ser Pro Ala Leu Ser Cys Asp Arg Gly His Arg Val Ser  
370 375 380

<210> 34  
<211> 1224  
<212> PRT  
<213> Homo sapiens

<400> 34

Met Glu Pro Leu Leu Leu Gly Arg Gly Leu Ile Val Tyr Leu Met Phe  
1 5 10 15

Leu Leu Leu Lys Phe Ser Lys Ala Ile Glu Ile Pro Ser Ser Val Gln  
20 25 30

Gln Val Pro Thr Ile Ile Lys Gln Ser Lys Val Gln Val Ala Phe Pro  
35 40 45

Phe Asp Glu Tyr Phe Gln Ile Glu Cys Glu Ala Lys Gly Asn Pro Glu  
50 55 60

Pro Thr Phe Ser Trp Thr Lys Asp Gly Asn Pro Phe Tyr Phe Thr Asp  
65 70 75 80

His Arg Ile Ile Pro Ser Asn Asn Ser Gly Thr Phe Arg Ile Pro Asn  
85 90 95

Glu Gly His Ile Ser His Phe Gln Gly Lys Tyr Arg Cys Phe Ala Ser  
100 105 110

Asn Lys Leu Gly Ile Ala Met Ser Glu Glu Ile Glu Phe Ile Val Pro  
115 120 125

Ser Val Pro Lys Phe Pro Lys Glu Lys Ile Asp Pro Leu Glu Val Glu  
130 135 140

Glu Gly Asp Pro Ile Val Leu Pro Cys Asn Pro Pro Lys Gly Leu Pro  
145 150 155 160

Pro Leu His Ile Tyr Trp Met Asn Ile Glu Leu Glu His Ile Glu Gln  
165 170 175

Asp Glu Arg Val Tyr Met Ser Gln Lys Gly Asp Leu Tyr Phe Ala Asn  
180 185 190

Val Glu Glu Lys Asp Ser Arg Asn Asp Tyr Cys Cys Phe Ala Ala Phe  
195 200 205

Pro Arg Leu Arg Thr Ile Val Gln Lys Met Pro Met Lys Leu Thr Val  
210 215 220

Asn Ser Leu Lys His Ala Asn Asp Ser Ser Ser Thr Glu Ile Gly  
225 230 235 240

Ser Lys Ala Asn Ser Ile Lys Gln Arg Lys Pro Lys Leu Leu Pro  
245 250 255

Pro Thr Glu Ser Gly Ser Glu Ser Ser Ile Thr Ile Leu Lys Gly Glu  
260 265 270

Ile Leu Leu Leu Glu Cys Phe Ala Glu Gly Leu Pro Thr Pro Gln Val  
275 280 285

Asp Trp Asn Lys Ile Gly Gly Asp Leu Pro Lys Gly Arg Glu Ala Lys  
290 295 300

Glu Asn Tyr Gly Lys Thr Leu Lys Ile Glu Asn Val Ser Tyr Gln Asp  
305 310 315 320

Lys Gly Asn Tyr Arg Cys Thr Ala Ser Asn Phe Leu Gly Thr Ala Thr  
325 330 335

His Asp Phe His Val Ile Val Glu Glu Pro Pro Arg Trp Thr Lys Lys  
340 345 350

Pro Gln Ser Ala Val Tyr Ser Thr Gly Ser Asn Gly Ile Leu Leu Cys  
355 360 365

Glu Ala Glu Gly Glu Pro Gln Pro Thr Ile Lys Trp Arg Val Asn Gly  
370 375 380

Ser Pro Val Asp Asn His Pro Phe Ala Gly Asp Val Val Phe Pro Arg  
385 390 395 400

Glu Ile Ser Phe Thr Asn Leu Gln Pro Asn His Thr Ala Val Tyr Gln  
405 410 415

Cys Glu Ala Ser Asn Val His Gly Thr Ile Leu Ala Asn Ala Asn Ile  
420 425 430

Asp Val Val Asp Val Arg Pro Leu Ile Gln Thr Lys Asp Gly Glu Asn  
435 440 445

Tyr Ala Thr Val Val Gly Tyr Ser Ala Phe Leu His Cys Glu Phe Phe  
450 455 460

Ala Ser Pro Glu Ala Val Val Ser Trp Gln Lys Val Glu Glu Val Lys  
465 470 475 480

Pro Leu Glu Gly Arg Arg Tyr His Ile Tyr Glu Asn Gly Thr Leu Gln  
485 490 495

Ile Asn Arg Thr Thr Glu Glu Asp Ala Gly Ser Tyr Ser Cys Trp Val  
500 505 510

Glu Asn Ala Ile Gly Lys Thr Ala Val Thr Ala Asn Leu Asp Ile Arg  
515 520 525

Asn Ala Thr Lys Leu Arg Val Ser Pro Lys Asn Pro Arg Ile Pro Lys  
530 535 540

Leu His Met Leu Glu Leu His Cys Glu Ser Lys Cys Asp Ser His Leu  
545 550 555 560

Lys His Ser Leu Lys Leu Ser Trp Ser Lys Asp Gly Glu Ala Phe Glu  
565 570 575

Ile Asn Gly Thr Glu Asp Gly Arg Ile Ile Ile Asp Gly Ala Asn Leu  
580 585 590

Thr Ile Ser Asn Val Thr Leu Glu Asp Gln Gly Ile Tyr Cys Cys Ser  
595 600 605

Ala His Thr Ala Leu Asp Ser Ala Ala Asp Ile Thr Gln Val Thr Val  
610 615 620

Leu Asp Val Pro Asp Pro Pro Glu Asn Leu His Leu Ser Glu Arg Gln  
625 630 635 640

Asn Arg Ser Val Arg Leu Thr Trp Glu Ala Gly Ala Asp His Asn Ser  
645 650 655

Asn Ile Ser Glu Tyr Ile Val Glu Phe Glu Gly Asn Lys Glu Glu Pro  
660 665 670

Gly Arg Trp Glu Glu Leu Thr Arg Val Gln Gly Lys Lys Thr Thr Val  
675 680 685

Ile Leu Pro Leu Ala Pro Phe Val Arg Tyr Gln Phe Arg Val Ile Ala  
690 695 700

Val Asn Glu Val Gly Arg Ser Gln Pro Ser Gln Pro Ser Asp His His  
705 710 715 720

Glu Thr Pro Pro Ala Ala Pro Asp Arg Asn Pro Gln Asn Ile Arg Val  
725 730 735

Gln Ala Ser Gln Pro Lys Glu Met Ile Ile Lys Trp Glu Pro Leu Lys  
740 745 750

Ser Met Glu Gln Asn Gly Pro Gly Leu Glu Tyr Arg Val Thr Trp Lys  
755 760 765

Pro Gln Gly Ala Pro Val Glu Trp Glu Glu Thr Val Thr Asn His  
770 775 780

Thr Leu Arg Val Met Thr Pro Ala Val Tyr Ala Pro Tyr Asp Val Lys  
785 790 795 800

Val Gln Ala Ile Asn Gln Leu Gly Ser Gly Pro Asp Pro Gln Ser Val  
805 810 815

Thr Leu Tyr Ser Gly Glu Asp Tyr Pro Asp Thr Ala Pro Val Ile His  
820 825 830

Gly Val Asp Val Ile Asn Ser Thr Leu Val Lys Val Thr Trp Ser Thr  
835 840 845

Val Pro Lys Asp Arg Val His Gly Arg Leu Lys Gly Tyr Gln Ile Asn  
850 855 860

Trp Trp Lys Thr Lys Ser Leu Leu Asp Gly Arg Thr His Pro Lys Glu  
865 870 875 880

Val Asn Ile Leu Arg Phe Ser Gly Gln Arg Asn Ser Gly Met Val Pro  
885 890 895

Ser Leu Asp Ala Phe Ser Glu Phe His Leu Thr Val Leu Ala Tyr Asn  
900 905 910

Ser Lys Gly Ala Gly Pro Glu Ser Glu Pro Tyr Ile Phe Gln Thr Pro  
915 920 925

Glu Gly Val Pro Glu Gln Pro Thr Phe Leu Lys Val Ile Lys Val Asp  
930 935 940

Lys Asp Thr Ala Thr Leu Ser Trp Gly Leu Pro Lys Lys Leu Asn Gly  
945 950 955 960

Asn Leu Thr Gly Tyr Leu Leu Gln Tyr Gln Ile Ile Asn Asp Thr Tyr  
965 970 975

Glu Ile Gly Glu Leu Asn Asp Ile Asn Ile Thr Thr Pro Ser Lys Pro  
980 985 990

Ser Trp His Leu Ser Asn Leu Asn Ala Thr Thr Lys Tyr Lys Phe Tyr  
995 1000 1005

Leu Arg Ala Cys Thr Ser Gln Gly Cys Gly Lys Pro Ile Thr Glu  
1010 1015 1020

Glu Ser Ser Thr Leu Gly Glu Gly Ser Lys Gly Ile Gly Lys Ile  
1025 1030 1035

Ser Gly Val Asn Leu Thr Gln Lys Thr His Pro Val Glu Val Phe  
1040 1045 1050

Glu Pro Gly Ala Glu His Ile Val Arg Leu Met Thr Lys Asn Trp  
1055 1060 1065

Gly Asp Asn Asp Ser Ile Phe Gln Asp Val Ile Glu Thr Arg Gly  
1070 1075 1080

Arg Glu Tyr Ala Gly Leu Tyr Asp Asp Ile Ser Thr Gln Gly Trp  
1085 1090 1095

Phe Ile Gly Leu Met Cys Ala Ile Ala Leu Leu Thr Leu Leu Leu  
1100 1105 1110

Leu Thr Val Cys Phe Val Lys Arg Asn Arg Gly Gly Lys Tyr Ser  
1115 1120 1125

Val Lys Glu Lys Glu Asp Leu His Pro Asp Pro Glu Ile Gln Ser  
1130 1135 1140

Val Lys Asp Glu Thr Phe Gly Glu Tyr Ser Asp Ser Asp Glu Lys  
1145 1150 1155

Pro Leu Lys Gly Ser Leu Arg Ser Leu Asn Arg Asp Met Gln Pro  
1160 1165 1170

Thr Glu Ser Ala Asp Ser Leu Val Glu Tyr Gly Glu Gly Asp His  
1175 1180 1185

Gly Leu Phe Ser Glu Asp Gly Ser Phe Ile Gly Ala Tyr Ala Gly  
1190 1195 1200

Ser Lys Glu Lys Gly Ser Val Glu Ser Asn Gly Ser Ser Thr Ala  
1205 1210 1215

Thr Phe Pro Leu Arg Ala  
1220

<210> 35  
<211> 1253  
<212> PRT  
<213> Homo sapiens

<400> 35

Met Val Val Ala Leu Arg Tyr Val Trp Pro Leu Leu Cys Ser Pro  
1 5 10 15

Cys Leu Leu Ile Gln Ile Pro Glu Glu Tyr Glu Gly His His Val Met  
20 25 30

Glu Pro Pro Val Ile Thr Glu Gln Ser Pro Arg Arg Leu Val Val Phe  
35 40 45

Pro Thr Asp Asp Ile Ser Leu Lys Cys Glu Ala Ser Gly Lys Pro Glu  
50 55 60

Val Gln Phe Arg Trp Thr Arg Asp Gly Val His Phe Lys Pro Lys Glu  
65 70 75 80

Glu Leu Gly Val Thr Val Tyr Gln Ser Pro His Ser Gly Ser Phe Thr  
85 90 95

Ile Thr Gly Asn Asn Ser Asn Phe Ala Gln Arg Phe Gln Gly Ile Tyr  
100 105 110

Arg Cys Phe Ala Ser Asn Lys Leu Gly Thr Ala Met Ser His Glu Ile

115

120

125

Arg Leu Met Ala Glu Gly Ala Pro Lys Trp Pro Lys Glu Thr Val Lys  
 130 135 140

Pro Val Glu Val Glu Glu Gly Ser Val Val Leu Pro Cys Asn Pro  
 145 150 155 160

Pro Pro Ser Ala Glu Pro Leu Arg Ile Tyr Trp Met Asn Ser Lys Ile  
 165 170 175

Leu His Ile Lys Gln Asp Glu Arg Val Thr Met Gly Gln Asn Gly Asn  
 180 185 190

Leu Tyr Phe Ala Asn Val Leu Thr Ser Asp Asn His Ser Asp Tyr Ile  
 195 200 205

Cys His Ala His Phe Pro Gly Thr Arg Thr Ile Ile Gln Lys Glu Pro  
 210 215 220

Ile Asp Leu Arg Val Lys Ala Thr Asn Ser Met Ile Asp Arg Lys Pro  
 225 230 235 240

Arg Leu Leu Phe Pro Thr Asn Ser Ser His Leu Val Ala Leu Gln  
 245 250 255

Gly Gln Pro Leu Val Leu Glu Cys Ile Ala Glu Gly Phe Pro Thr Pro  
 260 265 270

Thr Ile Lys Trp Leu Arg Pro Ser Gly Pro Met Pro Ala Asp Arg Val  
 275 280 285

Thr Tyr Gln Asn His Asn Lys Thr Leu Gln Leu Leu Lys Val Gly Glu  
 290 295 300

Glu Asp Asp Gly Glu Tyr Arg Cys Leu Ala Glu Asn Ser Leu Gly Ser  
 305 310 315 320

Ala Arg His Ala Tyr Tyr Val Thr Val Glu Ala Ala Pro Tyr Trp Leu  
 325 330 335

His Lys Pro Gln Ser His Leu Tyr Gly Pro Gly Glu Thr Ala Arg Leu  
 340 345 350

Asp Cys Gln Val Gln Gly Arg Pro Gln Pro Glu Val Thr Trp Arg Ile  
 355 360 365

Asn Gly Ile Pro Val Glu Glu Leu Ala Lys Asp Gln Lys Tyr Arg Ile  
370 375 380

Gln Arg Gly Ala Leu Ile Leu Ser Asn Val Gln Pro Ser Asp Thr Met  
385 390 395 400

Val Thr Gln Cys Glu Ala Arg Asn Arg His Gly Leu Leu Leu Ala Asn  
405 410 415

Ala Tyr Ile Tyr Val Val Gln Leu Pro Ala Lys Ile Leu Thr Ala Asp  
420 425 430

Asn Gln Thr Tyr Met Ala Val Gln Gly Ser Thr Ala Tyr Leu Leu Cys  
435 440 445

Lys Ala Phe Gly Ala Pro Val Pro Ser Val Gln Trp Leu Asp Glu Asp  
450 455 460

Gly Thr Thr Val Leu Gln Asp Glu Arg Phe Phe Pro Tyr Ala Asn Gly  
465 470 475 480

Thr Leu Gly Ile Arg Asp Leu Gln Ala Asn Asp Thr Gly Arg Tyr Phe  
485 490 495

Cys Leu Ala Ala Asn Asp Gln Asn Asn Val Thr Ile Met Ala Asn Leu  
500 505 510

Lys Val Lys Asp Ala Thr Gln Ile Thr Gln Gly Pro Arg Ser Thr Ile  
515 520 525

Glu Lys Lys Gly Ser Arg Val Thr Phe Thr Cys Gln Ala Ser Phe Asp  
530 535 540

Pro Ser Leu Gln Pro Ser Ile Thr Trp Arg Gly Asp Gly Arg Asp Leu  
545 550 555 560

Gln Glu Leu Gly Asp Ser Asp Lys Tyr Phe Ile Glu Asp Gly Arg Leu  
565 570 575

Val Ile His Ser Leu Asp Tyr Ser Asp Gln Gly Asn Tyr Ser Cys Val  
580 585 590

Ala Ser Thr Glu Leu Asp Val Val Glu Ser Arg Ala Gln Leu Leu Val  
595 600 605

Val Gly Ser Pro Gly Pro Val Pro Arg Leu Val Leu Ser Asp Leu His

610

615

620

Leu Leu Thr Gln Ser Gln Val Arg Val Ser Trp Ser Pro Ala Glu Asp  
 625                           630                           635                           640

His Asn Ala Pro Ile Glu Lys Tyr Asp Ile Glu Phe Glu Asp Lys Glu  
 645                           650                           655

Met Ala Pro Glu Lys Trp Tyr Ser Leu Gly Lys Val Pro Gly Asn Gln  
 660                           665                           670

Thr Ser Thr Thr Leu Lys Leu Ser Pro Tyr Val His Tyr Thr Phe Arg  
 675                           680                           685

Val Thr Ala Ile Asn Lys Tyr Gly Pro Gly Glu Pro Ser Pro Val Ser  
 690                           695                           700

Glu Thr Val Val Thr Pro Glu Ala Ala Pro Glu Lys Asn Pro Val Asp  
 705                           710                           715                           720

Val Lys Gly Glu Gly Asn Glu Thr Thr Asn Met Val Ile Thr Trp Lys  
 725                           730                           735

Pro Leu Arg Trp Met Asp Trp Asn Ala Pro Gln Val Gln Tyr Arg Val  
 740                           745                           750

Gln Trp Arg Pro Gln Gly Thr Arg Gly Pro Trp Gln Glu Gln Ile Val  
 755                           760                           765

Ser Asp Pro Phe Leu Val Val Ser Asn Thr Ser Thr Phe Val Pro Tyr  
 770                           775                           780

Glu Ile Lys Val Gln Ala Val Asn Ser Gln Gly Lys Gly Pro Glu Pro  
 785                           790                           795                           800

Gln Val Thr Ile Gly Tyr Ser Gly Glu Asp Tyr Pro Gln Ala Ile Pro  
 805                           810                           815

Glu Leu Glu Gly Ile Glu Ile Leu Asn Ser Ser Ala Val Leu Val Lys  
 820                           825                           830

Trp Arg Pro Val Asp Leu Ala Gln Val Lys Gly His Leu Arg Gly Tyr  
 835                           840                           845

Asn Val Thr Tyr Trp Arg Glu Gly Ser Gln Arg Lys His Ser Lys Arg  
 850                           855                           860

His Ile His Lys Asp His Val Val Val Pro Ala Asn Thr Thr Ser Val  
865 870 875 880

Ile Leu Ser Gly Leu Arg Pro Tyr Ser Ser Tyr His Leu Glu Val Gln  
885 890 895

Ala Phe Asn Gly Arg Gly Ser Gly Pro Ala Ser Glu Phe Thr Phe Ser  
900 905 910

Thr Pro Glu Gly Val Pro Gly His Pro Glu Ala Leu His Leu Glu Cys  
915 920 925

Gln Ser Asn Thr Ser Leu Leu Leu Arg Trp Gln Pro Pro Leu Ser His  
930 935 940

Asn Gly Val Leu Thr Gly Tyr Val Leu Ser Tyr His Pro Leu Asp Glu  
945 950 955 960

Gly Gly Lys Gly Gln Leu Ser Phe Asn Leu Arg Asp Pro Glu Leu Arg  
965 970 975

Thr His Asn Leu Thr Asp Leu Ser Pro His Leu Arg Tyr Arg Phe Gln  
980 985 990

Leu Gln Ala Thr Thr Lys Glu Gly Pro Gly Glu Ala Ile Val Arg Glu  
995 1000 1005

Gly Gly Thr Met Ala Leu Ser Gly Ile Ser Asp Phe Gly Asn Ile  
1010 1015 1020

Ser Ala Thr Ala Gly Glu Asn Tyr Ser Val Val Ser Trp Val Pro  
1025 1030 1035

Lys Glu Gly Gln Cys Asn Phe Arg Phe His Ile Leu Phe Lys Ala  
1040 1045 1050

Leu Gly Glu Glu Lys Gly Gly Ala Ser Leu Ser Pro Gln Tyr Val  
1055 1060 1065

Ser Tyr Asn Gln Ser Ser Tyr Thr Gln Trp Asp Leu Gln Pro Asp  
1070 1075 1080

Thr Asp Tyr Glu Ile His Leu Phe Lys Glu Arg Met Phe Arg His  
1085 1090 1095

Gln Met Ala Val Lys Thr Asn Gly Thr Gly Arg Val Arg Leu Pro

1100

1105

1110

Pro Ala Gly Phe Ala Thr Glu Gly Trp Phe Ile Gly Phe Val Ser  
1115 1120 1125

Ala Ile Ile Leu Leu Leu Val Leu Leu Ile Leu Cys Phe Ile  
1130 1135 1140

Lys Arg Ser Lys Gly Gly Lys Tyr Ser Val Lys Asp Lys Glu Asp  
1145 1150 1155

Thr Gln Val Asp Ser Glu Ala Arg Pro Met Lys Asp Glu Thr Phe  
1160 1165 1170

Gly Glu Tyr Ser Asp Asn Glu Glu Lys Ala Phe Gly Ser Ser Gln  
1175 1180 1185

Pro Ser Leu Asn Gly Asp Ile Lys Pro Leu Gly Ser Asp Asp Ser  
1190 1195 1200

Leu Ala Asp Tyr Gly Gly Ser Val Asp Val Gln Phe Asn Glu Asp  
1205 1210 1215

Gly Ser Phe Ile Gly Gln Tyr Ser Gly Lys Lys Glu Lys Glu Ala  
1220 1225 1230

Ala Gly Gly Asn Asp Ser Ser Gly Ala Thr Ser Pro Ile Asn Pro  
1235 1240 1245

Ala Val Ala Leu Glu  
1250

<210> 36  
<211> 1066  
<212> PRT  
<213> Homo sapiens

<400> 36

Met Ala Arg Gln Pro Pro Pro Pro Trp Val His Ala Ala Phe Leu Leu  
1 5 10 15

Cys Leu Leu Ser Leu Gly Gly Ala Ile Glu Ile Pro Met Asp Pro Ser  
20 25 30

Ile Gln Asn Glu Leu Thr Gln Pro Pro Thr Ile Thr Lys Gln Ser Ala  
35 40 45

Lys Asp His Ile Val Asp Pro Arg Asp Asn Ile Leu Ile Glu Cys Glu  
50 55 60

Ala Lys Gly Asn Pro Ala Pro Ser Phe His Trp Thr Arg Asn Ser Arg  
65 70 75 80

Phe Phe Asn Ile Ala Lys Asp Pro Arg Val Ser Met Arg Arg Arg Ser  
85 90 95

Gly Thr Leu Val Ile Asp Phe Arg Ser Gly Gly Arg Pro Glu Glu Tyr  
100 105 110

Glu Gly Glu Tyr Gln Cys Phe Ala Arg Asn Lys Phe Gly Thr Ala Leu  
115 120 125

Ser Asn Arg Ile Arg Leu Gln Val Ser Lys Ser Pro Leu Trp Pro Lys  
130 135 140

Glu Asn Leu Asp Pro Val Val Val Gln Glu Gly Ala Pro Leu Thr Leu  
145 150 155 160

Gln Cys Asn Pro Pro Pro Gly Leu Pro Ser Pro Val Ile Phe Trp Met  
165 170 175

Ser Ser Ser Met Glu Pro Ile Thr Gln Asp Lys Arg Val Ser Gln Gly  
180 185 190

His Asn Gly Asp Leu Tyr Phe Ser Asn Val Met Leu Gln Asp Met Gln  
195 200 205

Thr Asp Tyr Ser Cys Asn Ala Arg Phe His Phe Thr His Thr Ile Gln  
210 215 220

Gln Lys Asn Pro Phe Thr Leu Lys Val Leu Thr Thr Arg Gly Val Ala  
225 230 235 240

Glu Arg Thr Pro Ser Phe Met Tyr Pro Gln Gly Thr Ala Ser Ser Gln  
245 250 255

Met Val Leu Arg Gly Met Asp Leu Leu Leu Glu Cys Ile Ala Ser Gly  
260 265 270

Val Pro Thr Pro Asp Ile Ala Trp Tyr Lys Lys Gly Gly Asp Leu Pro  
275 280 285

Ser Asp Lys Ala Lys Phe Glu Asn Phe Asn Lys Ala Leu Arg Ile Thr  
290 295 300

Asn Val Ser Glu Glu Asp Ser Gly Glu Tyr Phe Cys Leu Ala Ser Asn  
305 310 315 320

Lys Met Gly Ser Ile Arg His Thr Ile Ser Val Arg Val Lys Ala Ala  
325 330 335

Pro Tyr Trp Leu Asp Glu Pro Lys Asn Leu Ile Leu Ala Pro Gly Glu  
340 345 350

Asp Gly Arg Leu Val Cys Arg Ala Asn Gly Asn Pro Lys Pro Thr Val  
355 360 365

Gln Trp Met Val Asn Gly Glu Pro Leu Gln Ser Ala Pro Pro Asn Pro  
370 375 380

Asn Arg Glu Val Ala Gly Asp Thr Ile Ile Phe Arg Asp Thr Gln Ile  
385 390 395 400

Ser Ser Arg Ala Val Tyr Gln Cys Asn Thr Ser Asn Glu His Gly Tyr  
405 410 415

Leu Leu Ala Asn Ala Phe Val Ser Val Leu Asp Val Pro Pro Arg Met  
420 425 430

Leu Ser Pro Arg Asn Gln Leu Ile Arg Val Ile Leu Tyr Asn Arg Thr  
435 440 445

Arg Leu Asp Cys Pro Phe Phe Gly Ser Pro Ile Pro Thr Leu Arg Trp  
450 455 460

Phe Lys Asn Gly Gln Gly Ser Asn Leu Asp Gly Gly Asn Tyr His Val  
465 470 475 480

Tyr Glu Asn Gly Ser Leu Glu Ile Lys Met Ile Arg Lys Glu Asp Gln  
485 490 495

Gly Ile Tyr Thr Cys Val Ala Thr Asn Ile Leu Gly Lys Ala Glu Asn  
500 505 510

Gln Val Arg Leu Glu Val Lys Asp Pro Thr Arg Ile Tyr Arg Met Pro  
515 520 525

Glu Asp Gln Val Ala Arg Arg Gly Thr Thr Val Gln Leu Glu Cys Arg  
530 535 540

Val Lys His Asp Pro Ser Leu Lys Leu Thr Val Ser Trp Leu Lys Asp  
545 550 555 560

Asp Glu Pro Leu Tyr Ile Gly Asn Arg Met Lys Lys Glu Asp Asp Ser  
565 570 575

Leu Thr Ile Phe Gly Val Ala Glu Arg Asp Gln Gly Ser Tyr Thr Cys  
580 585 590

Val Ala Ser Thr Glu Leu Asp Gln Asp Leu Ala Lys Ala Tyr Leu Thr  
595 600 605

Val Leu Ala Asp Gln Ala Thr Pro Thr Asn Arg Leu Ala Ala Leu Pro  
610 615 620

Lys Gly Arg Pro Asp Arg Pro Arg Asp Leu Glu Leu Thr Asp Leu Ala  
625 630 635 640

Glu Arg Ser Val Arg Leu Thr Trp Ile Pro Gly Asp Ala Asn Asn Ser  
645 650 655

Pro Ile Thr Asp Tyr Val Val Gln Phe Glu Asp Gln Phe Gln Pro  
660 665 670

Gly Val Trp His Asp His Ser Lys Tyr Pro Gly Ser Val Asn Ser Ala  
675 680 685

Val Leu Arg Leu Ser Pro Tyr Val Asn Tyr Gln Phe Arg Val Ile Ala  
690 695 700

Ile Asn Glu Val Gly Ser Ser His Pro Ser Leu Pro Ser Glu Arg Tyr  
705 710 715 720

Arg Thr Ser Gly Ala Pro Pro Glu Ser Asn Pro Gly Asp Val Lys Gly  
725 730 735

Glu Gly Thr Arg Lys Asn Asn Met Glu Ile Thr Trp Thr Pro Met Asn  
740 745 750

Ala Thr Ser Ala Phe Gly Pro Asn Leu Arg Tyr Ile Val Lys Trp Arg  
755 760 765

Arg Arg Glu Thr Arg Glu Ala Trp Asn Asn Val Thr Val Trp Gly Ser  
770 775 780

Arg Tyr Val Val Gly Gln Thr Pro Val Tyr Val Pro Tyr Glu Ile Arg  
785 790 795 800

Val Gln Ala Glu Asn Asp Phe Gly Lys Gly Pro Glu Pro Glu Ser Val  
805 810 815

Ile Gly Tyr Ser Gly Glu Asp Leu Pro Ser Ala Pro Arg Arg Phe Arg  
820 825 830

Val Arg Gln Pro Asn Leu Glu Thr Ile Asn Leu Glu Trp Asp His Pro  
835 840 845

Glu His Pro Asn Gly Ile Met Ile Gly Tyr Thr Leu Lys Tyr Val Ala  
850 855 860

Phe Asn Gly Thr Lys Val Gly Lys Gln Ile Val Glu Asn Phe Ser Pro  
865 870 875 880

Asn Gln Thr Lys Phe Thr Val Gln Arg Thr Asp Pro Val Ser Arg Tyr  
885 890 895

Arg Phe Thr Leu Ser Ala Arg Thr Gln Val Gly Ser Gly Glu Ala Val  
900 905 910

Thr Glu Glu Ser Pro Ala Pro Pro Asn Glu Ala Tyr Thr Asn Asn Gln  
915 920 925

Ala Asp Ile Ala Thr Gln Gly Trp Phe Ile Gly Leu Met Cys Ala Ile  
930 935 940

Ala Leu Leu Val Leu Ile Leu Leu Ile Val Cys Phe Ile Lys Arg Ser  
945 950 955 960

Arg Gly Gly Lys Tyr Pro Val Arg Glu Lys Lys Asp Val Pro Leu Gly  
965 970 975

Pro Glu Asp Pro Lys Glu Glu Asp Gly Ser Phe Asp Tyr Ser Asp Glu  
980 985 990

Asp Asn Lys Pro Leu Gln Gly Ser Gln Thr Ser Leu Asp Gly Thr Ile  
995 1000 1005

Lys Gln Gln Glu Ser Asp Asp Ser Leu Val Asp Tyr Gly Glu Gly  
1010 1015 1020

Gly Glu Gly Gln Phe Asn Glu Asp Gly Ser Phe Ile Gly Gln Tyr  
1025 1030 1035

Thr Val Lys Lys Asp Lys Glu Glu Thr Glu Gly Asn Glu Ser Ser  
1040 1045 1050

Glu Ala Thr Ser Pro Val Asn Ala Ile Tyr Ser Leu Ala  
1055 1060 1065

<210> 37  
<211> 280  
<212> PRT  
<213> Homo sapiens

<400> 37

Met Lys Phe Arg Ala Lys Ile Val Asp Gly Ala Cys Leu Asn His Phe  
1 5 10 15

Thr Arg Ile Ser Asn Met Ile Ala Lys Leu Ala Lys Thr Cys Thr Leu  
20 25 30

Arg Ile Ser Pro Asp Lys Leu Asn Phe Ile Leu Cys Asp Lys Leu Ala  
35 40 45

Asn Gly Gly Val Ser Met Trp Cys Glu Leu Glu Gln Glu Asn Phe Phe  
50 55 60

Asn Glu Phe Gln Met Glu Gly Val Ser Ala Glu Asn Asn Glu Ile Tyr  
65 70 75 80

Leu Glu Leu Thr Ser Glu Asn Leu Ser Arg Ala Leu Lys Thr Ala Gln  
85 90 95

Asn Ala Arg Ala Leu Lys Ile Lys Leu Thr Asn Lys His Phe Pro Cys  
100 105 110

Leu Thr Val Ser Val Glu Leu Leu Ser Met Ser Ser Ser Arg Ile  
115 120 125

Val Thr His Asp Ile Pro Ile Lys Val Ile Pro Arg Lys Leu Trp Lys  
130 135 140

Asp Leu Gln Glu Pro Val Val Pro Asp Pro Asp Val Ser Ile Tyr Leu  
145 150 155 160

Pro Val Leu Lys Thr Met Lys Ser Val Val Glu Lys Met Lys Asn Ile  
165 170 175

Ser Asn His Leu Val Ile Glu Ala Asn Leu Asp Gly Glu Leu Asn Leu  
180 185 190

Lys Ile Glu Thr Glu Leu Val Cys Val Thr Thr His Phe Lys Asp Leu  
195 200 205

Gly Asn Pro Pro Leu Ala Ser Glu Ser Thr His Glu Asp Arg Asn Val  
210 215 220

Glu His Met Ala Glu Val His Ile Asp Ile Arg Lys Leu Leu Gln Phe  
225 230 235 240

Leu Ala Gly Gln Gln Val Asn Pro Thr Lys Ala Leu Cys Asn Ile Val  
245 250 255

Asn Asn Lys Met Val His Phe Asp Leu Leu His Glu Asp Val Ser Leu  
260 265 270

Gln Tyr Phe Ile Pro Ala Leu Ser  
275 280

<210> 38  
<211> 278  
<212> PRT  
<213> Homo sapiens

<400> 38

Met Lys Phe Arg Ala Lys Ile Thr Gly Lys Gly Cys Leu Glu Leu Phe  
1 5 10 15

Ile His Val Ser Gly Thr Val Ala Arg Leu Ala Lys Val Cys Val Leu  
20 25 30

Arg Val Arg Pro Asp Ser Leu Cys Phe Gly Pro Ala Gly Ser Gly Gly  
35 40 45

Leu His Glu Ala Arg Leu Trp Cys Glu Val Arg Gln Gly Ala Phe Gln  
50 55 60

Gln Phe Arg Met Glu Gly Val Ser Glu Asp Leu Asp Glu Ile His Leu  
65 70 75 80

Glu Leu Thr Ala Glu His Leu Ser Arg Ala Ala Arg Ser Ala Ala Gly  
85 90 95

Ala Ser Ser Leu Lys Leu Gln Leu Thr His Lys Arg Arg Pro Ser Leu  
100 105 110

Thr Val Ala Val Glu Leu Val Ser Ser Leu Gly Arg Ala Arg Ser Val  
115 120 125

Val His Asp Leu Pro Val Arg Val Leu Pro Arg Arg Val Trp Arg Asp  
130 135 140

Cys Leu Pro Pro Ser Leu Arg Ala Ser Asp Ala Ser Ile Arg Leu Pro  
145 150 155 160

Arg Trp Arg Thr Leu Arg Ser Ile Val Glu Arg Met Ala Asn Val Gly  
165 170 175

Ser His Val Leu Val Glu Ala Asn Leu Ser Gly Arg Met Thr Leu Ser  
180 185 190

Ile Glu Thr Glu Val Val Ser Ile Gln Ser Tyr Phe Lys Asn Leu Gly  
195 200 205

Asn Pro Pro Gln Ser Ala Val Gly Val Pro Glu Asn Arg Asp Leu Glu  
210 215 220

Ser Met Val Gln Val Arg Val Asp Asn Arg Lys Leu Leu Gln Phe Leu  
225 230 235 240

Glu Gly Gln Gln Ile His Pro Thr Thr Ala Leu Cys Asn Ile Trp Asp  
245 250 255

Asn Thr Leu Leu Gln Leu Val Leu Val Gln Glu Tyr Val Ser Leu Gln  
260 265 270

Tyr Phe Ile Pro Ala Leu  
275

<210> 39  
<211> 1844  
<212> PRT  
<213> Homo sapiens  
  
<400> 39

Met Val Gly Val Leu Ala Met Ala Ala Ala Ala Pro Pro Pro Val  
1 5 10 15

Lys Asp Cys Glu Ile Glu Pro Cys Lys Lys Arg Lys Lys Asp Asp Asp  
20 25 30

Thr Ser Thr Cys Lys Thr Ile Thr Lys Tyr Leu Ser Pro Leu Gly Lys  
35 40 45

Thr Arg Asp Arg Val Phe Ala Pro Pro Lys Pro Ser Asn Ile Leu Asp

50

55

60

Tyr Phe Arg Lys Thr Ser Pro Thr Asn Glu Lys Thr Gln Leu Gly Lys  
 65 70 75 80

Glu Cys Lys Ile Lys Ser Pro Glu Ser Val Pro Val Asp Ser Asn Lys  
 85 90 95

Asp Cys Thr Thr Pro Leu Glu Met Phe Ser Asn Val Glu Phe Lys Lys  
 100 105 110

Lys Arg Lys Arg Val Asn Leu Ser His Gln Leu Asn Asn Ile Lys Thr  
 115 120 125

Glu Asn Glu Ala Pro Ile Glu Ile Ser Ser Asp Asp Ser Lys Glu Asp  
 130 135 140

Tyr Ser Leu Asn Asn Asp Phe Val Glu Ser Ser Thr Ser Val Leu Arg  
 145 150 155 160

Tyr Lys Lys Gln Val Glu Val Leu Ala Glu Asn Ile Gln Asp Thr Lys  
 165 170 175

Ser Gln Pro Asn Thr Met Thr Ser Leu Gln Asn Ser Lys Val Asn  
 180 185 190

Pro Lys Gln Gly Thr Thr Lys Asn Asp Phe Lys Lys Leu Arg Lys Arg  
 195 200 205

Lys Cys Arg Asp Val Val Asp Leu Ser Glu Ser Leu Pro Leu Ala Glu  
 210 215 220

Glu Leu Asn Leu Leu Lys Lys Asp Gly Lys Asp Thr Lys Gln Met Glu  
 225 230 235 240

Asn Thr Thr Ser His Ala Asn Ser Arg Asp Asn Val Thr Glu Ala Ala  
 245 250 255

Gln Leu Asn Asp Ser Ile Ile Thr Val Ser Tyr Glu Glu Phe Leu Lys  
 260 265 270

Ser His Lys Glu Asn Lys Val Glu Glu Ile Pro Asp Ser Thr Met Ser  
 275 280 285

Ile Cys Val Pro Ser Glu Thr Val Asp Glu Ile Val Lys Ser Gly Tyr  
 290 295 300

Ile Ser Glu Ser Glu Asn Ser Glu Ile Ser Gln Gln Val Arg Phe Lys  
305 310 315 320

Thr Val Thr Val Leu Ala Gln Val His Pro Ile Pro Pro Lys Lys Thr  
325 330 335

Gly Lys Ile Pro Arg Ile Phe Leu Lys Gln Lys Gln Phe Glu Met Glu  
340 345 350

Asn Ser Leu Ser Asp Pro Glu Asn Glu Gln Thr Val Gln Lys Arg Lys  
355 360 365

Ser Asn Val Val Ile Gln Glu Glu Leu Glu Leu Ala Val Leu Glu  
370 375 380

Ala Gly Ser Ser Glu Ala Val Lys Pro Lys Cys Thr Leu Glu Glu Arg  
385 390 395 400

Gln Gln Phe Met Lys Ala Phe Arg Gln Pro Ala Ser Asp Ala Leu Lys  
405 410 415

Asn Gly Val Lys Lys Ser Ser Asp Lys Gln Lys Asp Leu Asn Glu Lys  
420 425 430

Cys Leu Tyr Glu Val Gly Arg Asp Asp Asn Ser Lys Lys Ile Met Glu  
435 440 445

Asn Ser Gly Ile Gln Met Val Ser Lys Asn Gly Asn Leu Gln Leu His  
450 455 460

Thr Asp Lys Gly Ser Phe Leu Lys Glu Lys Asn Lys Lys Leu Lys Lys  
465 470 475 480

Lys Asn Lys Lys Thr Leu Asp Thr Gly Ala Ile Pro Gly Lys Asn Arg  
485 490 495

Glu Gly Asn Thr Gln Lys Lys Glu Thr Thr Phe Phe Leu Lys Glu Lys  
500 505 510

Gln Tyr Gln Asn Arg Met Ser Leu Arg Gln Arg Lys Thr Glu Phe Phe  
515 520 525

Lys Ser Ser Thr Leu Phe Asn Asn Glu Ser Leu Val Tyr Glu Asp Ile  
530 535 540

Ala Asn Asp Asp Leu Leu Lys Val Ser Ser Leu Cys Asn Asn Asn Lys

545

550

555

560

Leu Ser Arg Lys Thr Ser Ile Pro Val Lys Asp Ile Lys Leu Thr Gln  
 565                           570                           575

Ser Lys Ala Glu Ser Glu Ala Ser Leu Leu Asn Val Ser Thr Pro Lys  
 580                           585                           590

Ser Thr Arg Arg Ser Gly Arg Ile Ser Ser Thr Pro Thr Thr Glu Thr  
 595                           600                           605

Ile Arg Gly Ile Asp Ser Asp Asp Val Gln Asp Asn Ser Gln Leu Lys  
 610                           615                           620

Ala Ser Thr Gln Lys Ala Ala Asn Leu Ser Glu Lys His Ser Leu Tyr  
 625                           630                           640

Thr Ala Glu Leu Ile Thr Val Pro Phe Asp Ser Glu Ser Pro Ile Arg  
 645                           650                           655

Met Lys Phe Thr Arg Ile Ser Thr Pro Lys Lys Ser Lys Lys Ser  
 660                           665                           670

Asn Lys Arg Ser Glu Lys Ser Glu Ala Thr Asp Gly Gly Phe Thr Ser  
 675                           680                           685

Gln Ile Arg Lys Ala Ser Asn Thr Ser Lys Asn Ile Ser Lys Ala Lys  
 690                           695                           700

Gln Leu Ile Glu Lys Ala Lys Ala Leu His Ile Ser Arg Ser Lys Val  
 705                           710                           715                           720

Thr Glu Glu Ile Ala Ile Pro Leu Arg Arg Ser Ser Arg His Gln Thr  
 725                           730                           735

Leu Pro Glu Arg Lys Lys Leu Ser Glu Thr Glu Asp Ser Val Ile Ile  
 740                           745                           750

Ile Asp Ser Ser Pro Thr Ala Leu Lys His Pro Glu Lys Asn Gln Lys  
 755                           760                           765

Lys Leu Gln Cys Leu Asn Asp Val Leu Gly Lys Lys Leu Asn Thr Ser  
 770                           775                           780

Thr Lys Asn Val Pro Gly Lys Met Lys Val Ala Pro Leu Phe Leu Val  
 785                           790                           795                           800

Arg Lys Ala Gln Lys Ala Ala Asp Pro Val Pro Ser Phe Asp Glu Ser  
805 810 815

Ser Gln Asp Thr Ser Glu Lys Ser Gln Asp Cys Asp Val Gln Cys Lys  
820 825 830

Ala Lys Arg Asp Phe Leu Met Ser Gly Leu Pro Asp Leu Leu Lys Arg  
835 840 845

Gln Ile Ala Lys Lys Ala Ala Leu Asp Val Tyr Asn Ala Val Ser  
850 855 860

Thr Ser Phe Gln Arg Val Val His Val Gln Gln Lys Asp Asp Gly Cys  
865 870 875 880

Cys Leu Trp His Leu Lys Pro Pro Ser Cys Pro Leu Leu Thr Lys Phe  
885 890 895

Lys Glu Leu Asn Thr Lys Val Ile Asp Leu Ser Lys Cys Gly Ile Ala  
900 905 910

Leu Gly Glu Phe Ser Thr Leu Asn Ser Lys Leu Lys Ser Gly Asn Ser  
915 920 925

Ala Ala Val Phe Met Arg Thr Arg Lys Glu Phe Thr Glu Glu Val Arg  
930 935 940

Asn Leu Leu Leu Glu Glu Ile Arg Trp Ser Asn Pro Glu Phe Ser Leu  
945 950 955 960

Lys Lys Tyr Phe Pro Leu Leu Lys Lys Gln Ile Glu His Gln Val  
965 970 975

Leu Ser Ser Glu Cys His Ser Lys Gln Glu Leu Glu Ala Asp Val Ser  
980 985 990

His Lys Glu Thr Lys Arg Lys Leu Val Glu Ala Glu Asn Ser Lys Ser  
995 1000 1005

Lys Arg Lys Lys Pro Asn Glu Tyr Ser Lys Asn Leu Glu Lys Thr  
1010 1015 1020

Asn Arg Lys Ser Glu Glu Leu Ser Lys Arg Asn Asn Ser Ser Gly  
1025 1030 1035

Ile Lys Leu Asp Ser Ser Lys Asp Ser Gly Thr Glu Asp Met Leu

1040

1045

1050

Trp Thr Glu Lys Tyr Gln Pro Gln Thr Ala Ser Glu Leu Ile Gly  
 1055 1060 1065

Asn Glu Leu Ala Ile Lys Lys Leu His Ser Trp Leu Lys Asp Trp  
 1070 1075 1080

Lys Arg Arg Ala Glu Leu Glu Glu Arg Gln Asn Leu Lys Gly Lys  
 1085 1090 1095

Arg Asp Glu Lys His Glu Asp Phe Ser Gly Gly Ile Asp Phe Lys  
 1100 1105 1110

Gly Ser Ser Asp Asp Glu Glu Glu Ser Arg Leu Cys Asn Thr Val  
 1115 1120 1125

Leu Ile Thr Gly Pro Thr Gly Val Gly Lys Thr Ala Ala Val Tyr  
 1130 1135 1140

Ala Cys Ala Gln Glu Leu Gly Phe Lys Ile Phe Glu Val Asn Ala  
 1145 1150 1155

Ser Ser Gln Arg Ser Gly Arg Gln Ile Leu Ser Gln Leu Lys Glu  
 1160 1165 1170

Ala Thr Gln Ser His Gln Val Asp Lys Gln Gly Val Asn Ser Gln  
 1175 1180 1185

Lys Pro Cys Phe Phe Asn Ser Tyr Tyr Ile Gly Lys Ser Pro Lys  
 1190 1195 1200

Lys Ile Ser Ser Pro Lys Lys Val Val Thr Ser Pro Arg Lys Val  
 1205 1210 1215

Pro Pro Pro Ser Pro Lys Ser Ser Gly Pro Lys Arg Ala Leu Pro  
 1220 1225 1230

Pro Lys Thr Leu Ala Asn Tyr Phe Lys Val Ser Pro Lys Pro Lys  
 1235 1240 1245

Asn Asn Glu Glu Ile Gly Met Leu Leu Glu Asn Asn Lys Gly Ile  
 1250 1255 1260

Lys Asn Ser Phe Glu Gln Lys Gln Ile Thr Gln Thr Lys Ser Thr  
 1265 1270 1275

Asn Ala Thr Asn Ser Asn Val Lys Asp Val Gly Ala Glu Glu Pro  
1280 1285 1290

Ser Arg Lys Asn Ala Thr Ser Leu Ile Leu Phe Glu Glu Val Asp  
1295 1300 1305

Val Ile Phe Asp Glu Asp Ala Gly Phe Leu Asn Ala Ile Lys Thr  
1310 1315 1320

Phe Met Ala Thr Thr Lys Arg Pro Val Ile Leu Thr Thr Ser Asp  
1325 1330 1335

Pro Thr Phe Ser Leu Met Phe Asp Gly Cys Phe Glu Glu Ile Lys  
1340 1345 1350

Phe Ser Thr Pro Ser Leu Leu Asn Val Ala Ser Tyr Leu Gln Met  
1355 1360 1365

Ile Cys Leu Thr Glu Asn Phe Arg Thr Asp Val Lys Asp Phe Val  
1370 1375 1380

Thr Leu Leu Thr Ala Asn Thr Cys Asp Ile Arg Lys Ser Ile Leu  
1385 1390 1395

Tyr Leu Gln Phe Trp Ile Arg Ser Gly Gly Val Leu Glu Glu  
1400 1405 1410

Arg Pro Leu Thr Leu Tyr Arg Gly Asn Ser Arg Asn Val Gln Leu  
1415 1420 1425

Val Cys Ser Glu His Gly Leu Asp Asn Lys Ile Tyr Pro Lys Asn  
1430 1435 1440

Thr Lys Lys Lys Arg Val Asp Leu Pro Lys Cys Asp Ser Gly Cys  
1445 1450 1455

Ala Glu Thr Leu Phe Gly Leu Lys Asn Ile Phe Ser Pro Ser Glu  
1460 1465 1470

Asp Leu Phe Ser Phe Leu Lys His Lys Ile Thr Met Lys Glu Glu  
1475 1480 1485

Trp His Lys Phe Ile Gln Leu Leu Thr Glu Phe Gln Met Arg Asn  
1490 1495 1500

Val Asp Phe Leu Tyr Ser Asn Leu Glu Phe Ile Leu Pro Leu Pro

1505

1510

1515

Val Asp Thr Ile Pro Glu Thr Lys Asn Phe Cys Gly Pro Ser Val  
 1520 1525 1530

Thr Val Asp Ala Ser Ala Ala Thr Lys Ser Met Asn Cys Leu Ala  
 1535 1540 1545

Arg Lys His Ser Glu Arg Glu Gln Pro Leu Lys Lys Ser Gln Lys  
 1550 1555 1560

Lys Lys Gln Lys Lys Thr Leu Val Ile Leu Asp Asp Ser Asp Leu  
 1565 1570 1575

Phe Asp Thr Asp Leu Asp Phe Pro Asp Gln Ser Ile Ser Leu Ser  
 1580 1585 1590

Ser Val Ser Ser Ser Asn Ala Glu Glu Ser Lys Thr Gly Asp  
 1595 1600 1605

Glu Glu Ser Lys Ala Arg Asp Lys Gly Asn Asn Pro Glu Thr Lys  
 1610 1615 1620

Lys Ser Ile Pro Cys Pro Pro Lys Thr Thr Ala Gly Lys Lys Cys  
 1625 1630 1635

Ser Ala Leu Val Ser His Cys Leu Asn Ser Leu Ser Glu Phe Met  
 1640 1645 1650

Asp Asn Met Ser Phe Leu Asp Ala Leu Leu Thr Asp Val Arg Glu  
 1655 1660 1665

Gln Asn Lys Tyr Gly Arg Asn Asp Phe Ser Trp Thr Asn Gly Lys  
 1670 1675 1680

Val Thr Ser Gly Leu Cys Asp Glu Phe Ser Leu Glu Ser Asn Asp  
 1685 1690 1695

Gly Trp Thr Ser Gln Ser Ser Gly Glu Leu Lys Ala Ala Ala Glu  
 1700 1705 1710

Ala Leu Ser Phe Thr Lys Cys Ser Ser Ala Ile Ser Lys Ala Leu  
 1715 1720 1725

Glu Thr Leu Asn Ser Cys Lys Lys Leu Gly Arg Asp Pro Thr Asn  
 1730 1735 1740

Asp Leu Thr Phe Tyr Val Ser Gln Lys Arg Asn Asn Val Tyr Phe  
1745 1750 1755

Ser Gln Ser Ala Ala Asn Leu Asp Asn Ala Trp Lys Arg Ile Ser  
1760 1765 1770

Val Ile Lys Ser Val Phe Ser Ser Arg Ser Leu Leu Tyr Val Gly  
1775 1780 1785

Asn Arg Gln Ala Ser Ile Ile Glu Tyr Leu Pro Thr Leu Arg Asn  
1790 1795 1800

Ile Cys Lys Thr Glu Lys Leu Lys Glu Gln Gly Lys Ser Lys Arg  
1805 1810 1815

Arg Phe Leu His Tyr Phe Glu Gly Ile His Leu Asp Ile Pro Lys  
1820 1825 1830

Glu Thr Val Asn Thr Leu Ala Ala Asp Phe Pro  
1835 1840

<210> 40  
<211> 1148  
<212> PRT  
<213> Homo sapiens

<400> 40

Met Asp Ile Arg Lys Phe Phe Gly Val Ile Pro Ser Gly Lys Leu  
1 5 10 15

Val Ser Glu Thr Val Lys Lys Asn Glu Lys Thr Lys Ser Asp Glu Glu  
20 25 30

Thr Leu Lys Ala Lys Lys Gly Ile Lys Glu Ile Lys Val Asn Ser Ser  
35 40 45

Arg Lys Glu Asp Asp Phe Lys Gln Lys Gln Pro Ser Lys Lys Arg  
50 55 60

Ile Ile Tyr Asp Ser Asp Ser Glu Ser Glu Glu Thr Leu Gln Val Lys  
65 70 75 80

Asn Ala Lys Lys Pro Pro Glu Lys Leu Pro Val Ser Ser Lys Pro Gly  
85 90 95

Lys Ile Ser Arg Gln Asp Pro Val Thr Tyr Ile Ser Glu Thr Asp Glu  
100 105 110

Glu Asp Asp Phe Met Cys Lys Lys Ala Ala Ser Lys Ser Lys Glu Asn  
115 120 125

Gly Arg Ser Thr Asn Ser His Leu Gly Thr Ser Asn Met Lys Lys Asn  
130 135 140

Glu Glu Asn Thr Lys Thr Lys Asn Lys Pro Leu Ser Pro Ile Lys Leu  
145 150 155 160

Thr Pro Thr Ser Val Leu Asp Tyr Phe Gly Thr Gly Ser Val Gln Arg  
165 170 175

Ser Asn Lys Lys Met Val Ala Ser Lys Arg Lys Glu Leu Ser Gln Asn  
180 185 190

Thr Asp Glu Ser Gly Leu Asn Asp Glu Ala Ile Ala Lys Gln Leu Gln  
195 200 205

Leu Asp Glu Asp Ala Glu Leu Glu Arg Gln Leu His Glu Asp Glu Glu  
210 215 220

Phe Ala Arg Thr Leu Ala Met Leu Asp Glu Glu Pro Lys Thr Lys Lys  
225 230 235 240

Ala Arg Lys Asp Thr Glu Ala Gly Glu Thr Phe Ser Ser Val Gln Ala  
245 250 255

Asn Leu Ser Lys Ala Glu Lys His Lys Tyr Pro His Lys Val Lys Thr  
260 265 270

Ala Gln Val Ser Asp Glu Arg Lys Ser Tyr Ser Pro Arg Lys Gln Ser  
275 280 285

Lys Tyr Glu Ser Ser Lys Glu Ser Gln Gln His Ser Lys Ser Ser Ala  
290 295 300

Asp Lys Ile Gly Glu Val Ser Ser Pro Lys Ala Ser Ser Lys Leu Ala  
305 310 315 320

Ile Met Lys Arg Lys Glu Glu Ser Ser Tyr Lys Glu Ile Glu Pro Val  
325 330 335

Ala Ser Lys Arg Lys Glu Asn Ala Ile Lys Leu Lys Gly Glu Thr Lys  
340 345 350

Thr Pro Lys Lys Thr Lys Ser Ser Pro Ala Lys Lys Glu Ser Val Ser  
355 360 365

Pro Glu Asp Ser Glu Lys Lys Arg Thr Asn Tyr Gln Ala Tyr Arg Ser  
370 375 380

Tyr Leu Asn Arg Glu Gly Pro Lys Ala Leu Gly Ser Lys Glu Ile Pro  
385 390 395 400

Lys Gly Ala Glu Asn Cys Leu Glu Gly Leu Ile Phe Val Ile Thr Gly  
405 410 415

Val Leu Glu Ser Ile Glu Arg Asp Glu Ala Lys Ser Leu Ile Glu Arg  
420 425 430

Tyr Gly Gly Lys Val Thr Gly Asn Val Ser Lys Lys Thr Asn Tyr Leu  
435 440 445

Val Met Gly Arg Asp Ser Gly Gln Ser Lys Ser Asp Lys Ala Ala Ala  
450 455 460

Leu Gly Thr Lys Ile Ile Asp Glu Asp Gly Leu Leu Asn Leu Ile Arg  
465 470 475 480

Thr Met Pro Gly Lys Lys Ser Lys Tyr Glu Ile Ala Val Glu Thr Glu  
485 490 495

Met Lys Lys Glu Ser Lys Leu Glu Arg Thr Pro Gln Lys Asn Val Gln  
500 505 510

Gly Lys Arg Lys Ile Ser Pro Ser Lys Lys Glu Ser Glu Ser Lys Lys  
515 520 525

Ser Arg Pro Thr Ser Lys Arg Asp Ser Leu Ala Lys Thr Ile Lys Lys  
530 535 540

Glu Thr Asp Val Phe Trp Lys Ser Leu Asp Phe Lys Glu Gln Val Ala  
545 550 555 560

Glu Glu Thr Ser Gly Asp Ser Lys Ala Arg Asn Leu Ala Asp Asp Ser  
565 570 575

Ser Glu Asn Lys Val Glu Asn Leu Leu Trp Val Asp Lys Tyr Lys Pro  
580 585 590

; Thr Ser Leu Lys Thr Ile Ile Gly Gln Gln Gly Asp Gln Ser Cys Ala  
595 600 605

Asn Lys Leu Leu Arg Trp Leu Arg Asn Trp Gln Lys Ser Ser Ser Glu  
610 615 620

Asp Lys Lys His Ala Ala Lys Phe Gly Lys Phe Ser Gly Lys Asp Asp  
625 630 635 640

Gly Ser Ser Phe Lys Ala Ala Leu Leu Ser Gly Pro Pro Gly Val Gly  
645 650 655

Lys Thr Thr Thr Ala Ser Leu Val Cys Gln Glu Leu Gly Tyr Ser Tyr  
660 665 670

Val Glu Leu Asn Ala Ser Asp Thr Arg Ser Lys Ser Ser Leu Lys Ala  
675 680 685

Ile Val Ala Glu Ser Leu Asn Asn Thr Ser Ile Lys Gly Phe Tyr Ser  
690 695 700

Asn Gly Ala Ala Ser Ser Val Ser Thr Lys His Ala Leu Ile Met Asp  
705 710 715 720

Glu Val Asp Gly Met Ala Gly Asn Glu Asp Arg Gly Gly Ile Gln Glu  
725 730 735

Leu Ile Gly Leu Ile Lys His Thr Lys Ile Pro Ile Ile Cys Met Cys  
740 745 750

Asn Asp Arg Asn His Pro Lys Ile Arg Ser Leu Val His Tyr Cys Phe  
755 760 765

Asp Leu Arg Phe Gln Arg Pro Arg Val Glu Gln Ile Lys Gly Ala Met  
770 775 780

Met Ser Ile Ala Phe Lys Glu Gly Leu Lys Ile Pro Pro Pro Ala Met  
785 790 795 800

Asn Glu Ile Ile Leu Gly Ala Asn Gln Asp Ile Arg Gln Val Leu His  
805 810 815

Asn Leu Ser Met Trp Cys Ala Arg Ser Lys Ala Leu Thr Tyr Asp Gln  
820 825 830

Ala Lys Ala Asp Ser His Arg Ala Lys Lys Asp Ile Lys Met Gly Pro  
835 840 845

Phe Asp Val Ala Arg Lys Val Phe Ala Ala Gly Glu Glu Thr Ala His  
850 855 860

Met Ser Leu Val Asp Lys Ser Asp Leu Phe Phe His Asp Tyr Ser Ile  
865 870 875 880

Ala Pro Leu Phe Val Gln Glu Asn Tyr Ile His Val Lys Pro Val Ala  
885 890 895

Ala Gly Gly Asp Met Lys Lys His Leu Met Leu Leu Ser Arg Ala Ala  
900 905 910

Asp Ser Ile Cys Asp Gly Asp Leu Val Asp Ser Gln Ile Arg Ser Lys  
915 920 925

Gln Asn Trp Ser Leu Leu Pro Ala Gln Ala Ile Tyr Ala Ser Val Leu  
930 935 940

Pro Gly Glu Leu Met Arg Gly Tyr Met Thr Gln Phe Pro Thr Phe Pro  
945 950 955 960

Ser Trp Leu Gly Lys His Ser Ser Thr Gly Lys His Asp Arg Ile Val  
965 970 975

Gln Asp Leu Ala Leu His Met Ser Leu Arg Thr Tyr Ser Ser Lys Arg  
980 985 990

Thr Val Asn Met Asp Tyr Leu Ser Leu Leu Arg Asp Ala Leu Val Gln  
995 1000 1005

Pro Leu Thr Ser Gln Gly Val Asp Gly Val Gln Asp Val Val Ala  
1010 1015 1020

Leu Met Asp Thr Tyr Tyr Leu Met Lys Glu Asp Phe Glu Asn Ile  
1025 1030 1035

Met Glu Ile Ser Ser Trp Gly Gly Lys Pro Ser Pro Phe Ser Lys  
1040 1045 1050

Leu Asp Pro Lys Val Lys Ala Ala Phe Thr Arg Ala Tyr Asn Lys  
1055 1060 1065

Glu Ala His Leu Thr Pro Tyr Ser Leu Gln Ala Ile Lys Ala Ser  
1070 1075 1080

Arg His Ser Thr Ser Pro Ser Leu Asp Ser Glu Tyr Asn Glu Glu  
1085 1090 1095

Leu Asn Glu Asp Asp Ser Gln Ser Asp Glu Lys Asp Gln Asp Ala  
1100 1105 1110

Ile Glu Thr Asp Ala Met Ile Lys Lys Lys Thr Lys Ser Ser Lys  
1115 1120 1125

Pro Ser Lys Pro Glu Lys Asp Lys Glu Pro Arg Lys Gly Lys Gly  
1130 1135 1140

Lys Ser Ser Lys Lys  
1145

<210> 41  
<211> 307  
<212> PRT  
<213> Homo sapiens

<400> 41

Met Ala Glu Ile Ser Asp Leu Asp Arg Gln Ile Glu Gln Leu Arg Arg  
1 5 10 15

Cys Glu Leu Ile Lys Glu Ser Glu Val Lys Ala Leu Cys Ala Lys Ala  
20 25 30

Arg Glu Ile Leu Val Glu Glu Ser Asn Val Gln Arg Val Asp Ser Pro  
35 40 45

Val Thr Val Cys Gly Asp Ile His Gly Gln Phe Tyr Asp Leu Lys Glu  
50 55 60

Leu Phe Arg Val Gly Gly Asp Val Pro Glu Thr Asn Tyr Leu Phe Met  
65 70 75 80

Gly Asp Phe Val Asp Arg Gly Phe Tyr Ser Val Glu Thr Phe Leu Leu  
85 90 95

Leu Leu Ala Leu Lys Val Arg Tyr Pro Asp Arg Ile Thr Leu Ile Arg  
100 105 110

Gly Asn His Glu Ser Arg Gln Ile Thr Gln Val Tyr Gly Phe Tyr Asp  
115 120 125

Glu Cys Leu Arg Lys Tyr Gly Ser Val Thr Val Trp Arg Tyr Cys Thr  
130 135 140

Glu Ile Phe Asp Tyr Leu Ser Leu Ser Ala Ile Ile Asp Gly Lys Ile

145 150 155 160

Phe Cys Val His Gly Gly Leu Ser Pro Ser Ile Gln Thr Leu Asp Gln  
165 170 175

Ile Arg Thr Ile Asp Arg Lys Gln Glu Val Pro His Asp Gly Pro Met  
180 185 190

Cys Asp Leu Leu Trp Ser Asp Pro Glu Asp Thr Thr Gly Trp Gly Val  
195 200 205

Ser Pro Arg Gly Ala Gly Tyr Leu Phe Gly Ser Asp Val Val Ala Gln  
210 215 220

Phe Asn Ala Ala Asn Asp Ile Asp Met Ile Cys Arg Ala His Gln Leu  
225 230 235 240

Val Met Glu Gly Tyr Lys Trp His Phe Asn Glu Thr Val Leu Thr Val  
245 250 255

Trp Ser Ala Pro Asn Tyr Cys Tyr Arg Cys Gly Asn Val Ala Ala Ile  
260 265 270

Leu Glu Leu Asp Glu His Leu Gln Lys Asp Phe Ile Ile Phe Glu Ala  
275 280 285

Ala Pro Gln Glu Thr Arg Gly Ile Pro Ser Lys Lys Pro Val Ala Asp  
290 295 300

Tyr Phe Leu  
305

<210> 42  
<211> 773  
<212> PRT  
<213> Homo sapiens

<400> 42

Met Phe Ser Leu Ser Ser Thr Val Gln Pro Gln Val Thr Val Pro Leu  
1 5 10 15

Ser His Leu Ile Asn Ala Phe His Thr Pro Lys Asn Thr Ser Val Ser  
20 25 30

Leu Ser Gly Val Ser Val Ser Gln Asn Gln His Arg Asp Val Val Pro  
35 40 45

Glu His Glu Ala Pro Ser Ser Glu Cys Met Phe Ser Asp Phe Leu Thr  
50 55 60

Lys Leu Asn Ile Val Ser Ile Gly Lys Gly Lys Ile Phe Glu Gly Tyr  
65 70 75 80

Arg Ser Met Phe Met Glu Pro Ala Lys Arg Met Lys Lys Ser Leu Asp  
85 90 95

Thr Thr Asp Asn Trp His Ile Arg Pro Glu Pro Phe Ser Leu Ser Ile  
100 105 110

Pro Pro Ser Leu Asn Leu Arg Asp Leu Gly Leu Ser Glu Leu Lys Ile  
115 120 125

Gly Gln Ile Asp Gln Leu Val Glu Asn Leu Leu Pro Gly Phe Cys Lys  
130 135 140

Gly Lys Asn Ile Ser Ser His Trp His Thr Ser His Val Ser Ala Gln  
145 150 155 160

Ser Phe Phe Glu Asn Lys Tyr Gly Asn Leu Asp Ile Phe Ser Thr Leu  
165 170 175

Arg Ser Ser Cys Leu Tyr Arg His His Ser Arg Ala Leu Gln Ser Ile  
180 185 190

Cys Ser Asp Leu Gln Tyr Trp Pro Val Phe Ile Gln Ser Arg Gly Phe  
195 200 205

Lys Thr Leu Lys Ser Arg Thr Arg Arg Leu Gln Ser Thr Ser Glu Arg  
210 215 220

Leu Ala Glu Thr Gln Asn Ile Ala Pro Ser Phe Val Lys Gly Phe Leu  
225 230 235 240

Leu Arg Asp Arg Gly Ser Asp Val Glu Ser Leu Asp Lys Leu Met Lys  
245 250 255

Thr Lys Asn Ile Pro Glu Ala His Gln Asp Ala Phe Lys Thr Gly Phe  
260 265 270

Ala Glu Gly Phe Leu Lys Ala Gln Ala Leu Thr Gln Lys Thr Asn Asp  
275 280 285

Ser Leu Arg Arg Thr Arg Leu Ile Leu Phe Val Leu Leu Phe Gly  
290 295 300

Ile Tyr Gly Leu Leu Lys Asn Pro Phe Leu Ser Val Arg Phe Arg Thr  
305 310 315 320

Thr Thr Gly Leu Asp Ser Ala Val Asp Pro Val Gln Met Lys Asn Val  
325 330 335

Thr Phe Glu His Val Lys Gly Val Glu Glu Ala Lys Gln Glu Leu Gln  
340 345 350

Glu Val Val Glu Phe Leu Lys Asn Pro Gln Lys Phe Thr Ile Leu Gly  
355 360 365

Gly Lys Leu Pro Lys Gly Ile Leu Leu Val Gly Pro Pro Gly Thr Gly  
370 375 380

Lys Thr Leu Leu Ala Arg Ala Val Ala Gly Glu Ala Asp Val Pro Phe  
385 390 395 400

Tyr Tyr Ala Ser Gly Ser Glu Phe Asp Glu Met Phe Val Gly Val Gly  
405 410 415

Ala Ser Arg Ile Arg Asn Leu Phe Arg Glu Ala Lys Ala Asn Ala Pro  
420 425 430

Cys Val Ile Phe Ile Asp Glu Leu Asp Ser Val Gly Gly Lys Arg Ile  
435 440 445

Glu Ser Pro Met His Pro Tyr Ser Arg Gln Thr Ile Asn Gln Leu Leu  
450 455 460

Ala Glu Met Asp Gly Phe Lys Pro Asn Glu Gly Val Ile Ile Ile Gly  
465 470 475 480

Ala Thr Asn Phe Pro Glu Ala Leu Asp Asn Ala Leu Ile Arg Pro Gly  
485 490 495

Arg Phe Asp Met Gln Val Thr Val Pro Arg Pro Asp Val Lys Gly Arg  
500 505 510

Thr Glu Ile Leu Lys Trp Tyr Leu Asn Lys Ile Lys Phe Asp Gln Ser  
515 520 525

Val Asp Pro Glu Ile Ile Ala Arg Gly Thr Val Gly Phe Ser Gly Ala  
530 535 540

Glu Leu Glu Asn Leu Val Asn Gln Ala Ala Leu Lys Ala Ala Val Asp  
545 550 555 560

Gly Lys Glu Met Val Thr Met Lys Glu Leu Glu Phe Ser Lys Asp Lys  
565 570 575

Ile Leu Met Gly Pro Glu Arg Arg Ser Val Glu Ile Asp Asn Lys Asn  
580 585 590

Lys Thr Ile Thr Ala Tyr His Glu Ser Gly His Ala Ile Ile Ala Tyr  
595 600 605

Tyr Thr Lys Asp Ala Met Pro Ile Asn Lys Ala Thr Ile Met Pro Arg  
610 615 620

Gly Pro Thr Leu Gly His Val Ser Leu Leu Pro Glu Asn Asp Arg Trp  
625 630 635 640

Asn Glu Thr Arg Ala Gln Leu Leu Ala Gln Met Asp Val Ser Met Gly  
645 650 655

Gly Arg Val Ala Glu Glu Leu Ile Phe Gly Thr Asp His Ile Thr Thr  
660 665 670

Gly Ala Ser Ser Asp Phe Asp Asn Ala Thr Lys Ile Ala Lys Arg Met  
675 680 685

Val Thr Lys Phe Gly Met Ser Glu Lys Leu Gly Val Met Thr Tyr Ser  
690 695 700

Asp Thr Gly Lys Leu Ser Pro Glu Thr Gln Ser Ala Ile Glu Gln Glu  
705 710 715 720

Ile Arg Ile Leu Leu Arg Asp Ser Tyr Glu Arg Ala Lys His Ile Leu  
725 730 735

Lys Thr His Ala Lys Glu His Lys Asn Leu Ala Glu Ala Leu Leu Thr  
740 745 750

Tyr Glu Thr Leu Asp Ala Lys Glu Ile Gln Ile Val Leu Glu Gly Lys  
755 760 765

Lys Leu Glu Val Arg  
770

<210> 43  
<211> 534

<212> PRT

<213> Homo sapiens

<400> 43

Met Phe Ser Trp Val Ser Lys Asp Ala Arg Arg Lys Lys Glu Pro Glu  
1 5 10 15

Leu Phe Gln Thr Val Ala Glu Gly Leu Arg Gln Leu Tyr Ala Gln Lys  
20 25 30

Leu Leu Pro Leu Glu Glu His Tyr Arg Phe His Glu Phe His Ser Pro  
35 40 45

Ala Leu Glu Asp Ala Asp Phe Asp Asn Lys Pro Met Val Leu Leu Val  
50 55 60

Gly Gln Tyr Ser Thr Gly Lys Thr Thr Phe Ile Arg His Leu Ile Glu  
65 70 75 80

Gln Asp Phe Pro Gly Met Arg Ile Gly Pro Glu Pro Thr Thr Asp Ser  
85 90 95

Phe Ile Ala Val Met His Gly Pro Thr Glu Gly Val Val Pro Gly Asn  
100 105 110

Ala Leu Val Val Asp Pro Arg Arg Pro Phe Arg Lys Leu Asn Arg Phe  
115 120 125

Gly Asn Ala Phe Leu Asn Arg Phe Met Cys Ala Gln Leu Pro Asn Pro  
130 135 140

Val Leu Asp Ser Ile Ser Ile Ile Asp Thr Pro Gly Ile Leu Ser Gly  
145 150 155 160

Glu Lys Gln Arg Ile Ser Arg Gly Tyr Asp Phe Ala Ala Val Leu Glu  
165 170 175

Trp Phe Ala Asp Cys Trp Asp Arg Ile Ile Leu Leu Phe Asp Ala His  
180 185 190

Lys Gln Asp Ile Ser His Glu Phe Ser Glu Val Ile Lys Ala Leu Lys  
195 200 205

Asn His Glu Asp Lys Ile Arg Met Val Leu Asn Lys Ala Asp Gln Ile  
210 215 220

Glu Thr Gln Gln Leu Met Arg Val Tyr Gly Ala Leu Met Trp Ser Leu

225

230

235

240

Gly Lys Ile Ile Asn Thr Pro Glu Val Val Arg Val Tyr Ile Gly Ser  
 245 250 255

Phe Trp Ser His Pro Leu Leu Ile Pro Asp Asn Arg Lys Leu Phe Glu  
 260 265 270

Ala Glu Glu Gln Asp Leu Phe Lys Asp Ile Gln Ser Leu Pro Arg Asn  
 275 280 285

Ala Ala Leu Arg Lys Leu Asn Asp Leu Ile Lys Arg Ala Arg Leu Ala  
 290 295 300

Lys Val His Ala Tyr Ile Ile Ser Ser Leu Lys Lys Glu Met Pro Asn  
 305 310 315 320

Val Phe Gly Lys Glu Ser Lys Lys Glu Leu Val Asn Asn Leu Gly  
 325 330 335

Glu Ile Tyr Gln Lys Ile Glu Arg Glu His Gln Ile Ser Pro Gly Asp  
 340 345 350

Phe Pro Ser Leu Arg Lys Met Gln Glu Leu Leu Gln Thr Gln Asp Phe  
 355 360 365

Ser Lys Phe Gln Ala Leu Lys Pro Lys Leu Leu Asp Thr Val Asp Asp  
 370 375 380

Met Leu Ala Asn Asp Ile Ala Arg Leu Met Val Met Val Arg Gln Glu  
 385 390 395 400

Glu Ser Leu Met Pro Ser Gln Val Val Lys Gly Gly Ala Phe Asp Gly  
 405 410 415

Thr Met Asn Gly Pro Phe Gly His Gly Tyr Gly Glu Gly Ala Gly Glu  
 420 425 430

Gly Ile Asp Asp Val Glu Trp Val Val Gly Lys Asp Lys Pro Ser Tyr  
 435 440 445

Asp Glu Ile Phe Tyr Thr Leu Ser Pro Val Asn Gly Lys Ile Thr Gly  
 450 455 460

Ala Asn Val Lys Lys Glu Met Val Lys Ser Lys Leu Pro Asn Thr Glu  
 465 470 475 480

Leu Gly Lys Ile Trp Lys Leu Ala Asp Val Asp Lys Asp Gly Leu Leu  
485 490 495

Asp Asp Glu Glu Phe Ala Leu Ala Asn His Leu Ile Lys Val Lys Leu  
500 505 510

Glu Gly His Glu Leu Pro Ala Asp Leu Pro Pro His Leu Val Pro Pro  
515 520 525

Ser Lys Arg Arg His Glu  
530

<210> 44  
<211> 543  
<212> PRT  
<213> Homo sapiens

<400> 44

Met Phe Ser Trp Leu Lys Arg Gly Gly Ala Arg Gly Gln Gln Pro Glu  
1 5 10 15

Ala Ile Arg Thr Val Thr Ser Ala Leu Lys Glu Leu Tyr Arg Thr Lys  
20 25 30

Leu Leu Pro Leu Glu Glu His Tyr Arg Phe Gly Ala Phe His Ser Pro  
35 40 45

Ala Leu Glu Asp Ala Asp Phe Asp Gly Lys Pro Met Val Leu Val Ala  
50 55 60

Gly Gln Tyr Ser Thr Gly Lys Thr Ser Phe Ile Gln Tyr Leu Leu Glu  
65 70 75 80

Gln Glu Val Pro Gly Ser Arg Val Gly Pro Glu Pro Thr Thr Asp Cys  
85 90 95

Phe Val Ala Val Met His Gly Asp Thr Glu Gly Thr Val Pro Gly Asn  
100 105 110

Ala Leu Val Val Asp Pro Asp Lys Pro Phe Arg Lys Leu Asn Pro Phe  
115 120 125

Gly Asn Thr Phe Leu Asn Arg Phe Met Cys Ala Gln Leu Pro Asn Gln  
130 135 140

Val Leu Glu Ser Ile Ser Ile Ile Asp Thr Pro Gly Ile Leu Ser Gly  
145 150 155 160

Ala Lys Gln Arg Val Ser Arg Gly Tyr Asp Phe Pro Ala Val Leu Arg  
165 170 175

Trp Phe Ala Glu Arg Val Asp Leu Ile Ile Leu Leu Phe Asp Ala His  
180 185 190

Lys Leu Glu Ile Ser Asp Glu Phe Ser Glu Ala Ile Gly Ala Leu Arg  
195 200 205

Gly His Glu Asp Lys Ile Arg Val Val Leu Asn Lys Ala Asp Met Val  
210 215 220

Glu Thr Gln Gln Leu Met Arg Val Tyr Gly Ala Leu Met Trp Ala Leu  
225 230 235 240

Gly Lys Val Val Gly Thr Pro Glu Val Leu Arg Val Tyr Ile Gly Ser  
245 250 255

Phe Trp Ser Gln Pro Leu Leu Val Pro Asp Asn Arg Arg Leu Phe Glu  
260 265 270

Leu Glu Glu Gln Asp Leu Phe Arg Asp Ile Gln Gly Leu Pro Arg His  
275 280 285

Ala Ala Leu Arg Lys Leu Asn Asp Leu Val Lys Arg Ala Arg Leu Val  
290 295 300

Arg Val His Ala Tyr Ile Ile Ser Tyr Leu Lys Lys Glu Met Pro Ser  
305 310 315 320

Val Phe Gly Lys Glu Asn Lys Lys Gln Leu Ile Leu Lys Leu Pro  
325 330 335

Val Ile Phe Ala Lys Ile Gln Leu Glu His His Ile Ser Pro Gly Asp  
340 345 350

Phe Pro Asp Cys Gln Lys Met Gln Glu Leu Leu Met Ala His Asp Phe  
355 360 365

Thr Lys Phe His Ser Leu Lys Pro Lys Leu Leu Glu Ala Leu Asp Glu  
370 375 380

Met Leu Thr His Asp Ile Ala Lys Leu Met Pro Leu Leu Arg Gln Glu  
385 390 395 400

Glu Leu Glu Ser Thr Glu Val Gly Val Gln Gly Gly Ala Phe Glu Gly.  
405 410 415

Thr His Met Gly Pro Phe Val Glu Arg Gly Pro Asp Glu Ala Met Glu  
420 425 430

Asp Gly Glu Glu Gly Ser Asp Asp Glu Ala Glu Trp Val Val Thr Lys  
435 440 445

Asp Lys Ser Lys Tyr Asp Glu Ile Phe Tyr Asn Leu Ala Pro Ala Asp  
450 455 460

Gly Lys Leu Ser Gly Ser Lys Ala Lys Thr Trp Met Val Gly Thr Lys  
465 470 475 480

Leu Pro Asn Ser Val Leu Gly Arg Ile Trp Lys Leu Ser Asp Val Asp  
485 490 495

Arg Asp Gly Met Leu Asp Asp Glu Glu Phe Ala Leu Ala Ser His Leu  
500 505 510

Ile Glu Ala Lys Leu Glu Gly His Gly Leu Pro Ala Asn Leu Pro Arg  
515 520 525

Arg Leu Val Pro Pro Ser Lys Arg Arg His Lys Gly Ser Ala Glu  
530 535 540

<210> 45  
<211> 535  
<212> PRT  
<213> Homo sapiens

<400> 45

Met Phe Ser Trp Leu Gly Thr Asp Asp Arg Arg Arg Lys Asp Pro Glu  
1 5 10 15

Val Phe Gln Thr Val Ser Glu Gly Leu Lys Lys Leu Tyr Lys Ser Lys  
20 25 30

Leu Leu Pro Leu Glu Glu His Tyr Arg Phe His Glu Phe His Ser Pro  
35 40 45

Ala Leu Glu Asp Ala Asp Phe Asp Asn Lys Pro Met Val Leu Leu Val  
50 55 60

Gly Gln Tyr Ser Thr Gly Lys Thr Thr Phe Ile Arg Tyr Leu Leu Glu  
65 70 75 80

Gln Asp Phe Pro Gly Met Arg Ile Gly Pro Glu Pro Thr Thr Asp Ser  
85 90 95

Phe Ile Ala Val Met Gln Gly Asp Met Glu Gly Ile Ile Pro Gly Asn  
100 105 110

Ala Leu Val Val Asp Pro Lys Lys Pro Phe Arg Lys Leu Asn Ala Phe  
115 120 125

Gly Asn Ala Phe Leu Asn Arg Phe Val Cys Ala Gln Leu Pro Asn Pro  
130 135 140

Val Leu Glu Ser Ile Ser Val Ile Asp Thr Pro Gly Ile Leu Ser Gly  
145 150 155 160

Glu Lys Gln Arg Ile Ser Arg Gly Tyr Asp Phe Ala Ala Val Leu Glu  
165 170 175

Trp Phe Ala Glu Arg Val Asp Arg Ile Ile Leu Leu Phe Asp Ala His  
180 185 190

Lys Leu Asp Ile Ser Asp Glu Phe Ser Glu Val Ile Lys Ala Leu Lys  
195 200 205

Asn His Glu Asp Lys Met Arg Val Val Leu Asn Lys Ala Asp Gln Ile  
210 215 220

Glu Thr Gln Gln Leu Met Arg Val Tyr Gly Ala Leu Met Trp Ser Leu  
225 230 235 240

Gly Lys Ile Val Asn Thr Pro Glu Val Ile Arg Val Tyr Ile Gly Ser  
245 250 255

Phe Trp Ser His Pro Leu Leu Ile Pro Asp Asn Arg Lys Leu Phe Glu  
260 265 270

Ala Glu Glu Gln Asp Leu Phe Arg Asp Ile Gln Ser Leu Pro Arg Asn  
275 280 285

Ala Ala Leu Arg Lys Leu Asn Asp Leu Ile Lys Arg Ala Arg Leu Ala  
290 295 300

Lys Val His Ala Tyr Ile Ile Ser Ser Leu Lys Lys Glu Met Pro Ser  
305 310 315 320

Val Phe Gly Lys Asp Asn Lys Lys Lys Glu Leu Val Asn Asn Leu Ala

325

330

335

Glu Ile Tyr Gly Arg Ile Glu Arg Glu His Gln Ile Ser Pro Gly Asp  
 340 345 350

Phe Pro Asn Leu Lys Arg Met Gln Asp Gln Leu Gln Ala Gln Asp Phe  
 355 360 365

Ser Lys Phe Gln Pro Leu Lys Ser Lys Leu Leu Glu Val Val Asp Asp  
 370 375 380

Met Leu Ala His Asp Ile Ala Gln Leu Met Val Leu Val Arg Gln Glu  
 385 390 395 400

Glu Ser Gln Arg Pro Ile Gln Met Val Lys Gly Gly Ala Phe Glu Gly  
 405 410 415

Thr Leu His Gly Pro Phe Gly His Gly Tyr Gly Glu Gly Ala Gly Glu  
 420 425 430

Gly Ile Asp Asp Ala Glu Trp Val Val Ala Arg Asp Lys Pro Met Tyr  
 435 440 445

Asp Glu Ile Phe Tyr Thr Leu Ser Pro Val Asp Gly Lys Ile Thr Gly  
 450 455 460

Ala Asn Ala Lys Lys Glu Met Val Arg Ser Lys Leu Pro Asn Ser Val  
 465 470 475 480

Leu Gly Lys Ile Trp Lys Leu Ala Asp Ile Asp Lys Asp Gly Met Leu  
 485 490 495

Asp Asp Asp Glu Phe Ala Leu Ala Asn His Leu Ile Lys Val Lys Leu  
 500 505 510

Glu Gly His Glu Leu Pro Asn Glu Leu Pro Ala His Leu Leu Pro Pro  
 515 520 525

Ser Lys Arg Lys Val Ala Glu  
 530 535

<210> 46  
 <211> 541  
 <212> PRT  
 <213> Homo sapiens

<400> 46

Met Phe Ser Trp Met Gly Arg Gln Ala Gly Gly Arg Glu Arg Ala Gly  
1 5 10 15

Gly Ala Asp Ala Val Gln Thr Val Thr Gly Gly Leu Arg Ser Leu Tyr  
20 25 30

Leu Arg Lys Val Leu Pro Leu Glu Glu Ala Tyr Arg Phe His Glu Phe  
35 40 45

His Ser Pro Ala Leu Glu Asp Ala Asp Phe Glu Asn Lys Pro Met Ile  
50 55 60

Leu Leu Val Gly Gln Tyr Ser Thr Gly Lys Thr Thr Phe Ile Arg Tyr  
65 70 75 80

Leu Leu Glu Gln Asp Phe Pro Gly Met Arg Ile Gly Pro Glu Pro Thr  
85 90 95

Thr Asp Ser Phe Ile Ala Val Met Tyr Gly Glu Thr Glu Gly Ser Thr  
100 105 110

Pro Gly Asn Ala Leu Val Val Asp Pro Lys Lys Pro Phe Arg Lys Leu  
115 120 125

Ser Arg Phe Gly Asn Ala Phe Leu Asn Arg Phe Met Cys Ser Gln Leu  
130 135 140

Pro Asn Gln Val Leu Lys Ser Ile Ser Val Ile Asp Ser Pro Gly Ile  
145 150 155 160

Leu Ser Gly Glu Lys Gln Arg Ile Ser Arg Gly Tyr Asp Phe Cys Gln  
165 170 175

Val Leu Gln Trp Phe Ala Glu Arg Val Asp Arg Ile Ile Leu Leu Phe  
180 185 190

Asp Ala His Lys Leu Asp Ile Ser Asp Glu Phe Ser Glu Ala Ile Lys  
195 200 205

Ala Phe Arg Gly Gln Asp Asp Lys Ile Arg Val Val Leu Asn Lys Ala  
210 215 220

Asp Gln Val Asp Thr Gln Gln Leu Met Arg Val Tyr Gly Ala Leu Met  
225 230 235 240

Trp Ser Leu Gly Lys Val Ile Asn Thr Pro Glu Val Leu Arg Val Tyr  
245 250 255

Ile Gly Ser Phe Trp Ala Gln Pro Leu Gln Asn Thr Asp Asn Arg Arg  
260 265 270

Leu Phe Glu Ala Glu Ala Gln Asp Leu Phe Arg Asp Ile Gln Ser Leu  
275 280 285

Pro Gln Lys Ala Ala Val Arg Lys Leu Asn Asp Leu Ile Lys Arg Ala  
290 295 300

Arg Leu Ala Lys Val His Ala Tyr Ile Ile Ser Tyr Leu Lys Lys Glu  
305 310 315 320

Met Pro Ser Val Phe Gly Lys Glu Asn Lys Lys Arg Glu Leu Ile Ser  
325 330 335

Arg Leu Pro Glu Ile Tyr Ile Gln Leu Gln Arg Glu Tyr Gln Ile Ser  
340 345 350

Ala Gly Asp Phe Pro Glu Val Lys Ala Met Gln Glu Gln Leu Glu Asn  
355 360 365

Tyr Asp Phe Thr Lys Phe His Ser Leu Lys Pro Lys Leu Ile Glu Ala  
370 375 380

Val Asp Asn Met Leu Ser Asn Lys Ile Ser Pro Leu Met Asn Leu Ile  
385 390 395 400

Ser Gln Glu Glu Thr Ser Thr Pro Thr Gln Leu Val Gln Gly Gly Ala  
405 410 415

Phe Asp Gly Thr Thr Glu Gly Pro Phe Asn Gln Gly Tyr Gly Glu Gly  
420 425 430

Ala Lys Glu Gly Ala Asp Glu Glu Glu Trp Val Val Ala Lys Asp Lys  
435 440 445

Pro Val Tyr Asp Glu Leu Phe Tyr Thr Leu Ser Pro Ile Asn Gly Lys  
450 455 460

Ile Ser Gly Val Asn Ala Lys Lys Glu Met Val Thr Ser Lys Leu Pro  
465 470 475 480

Asn Ser Val Leu Gly Lys Ile Trp Lys Leu Ala Asp Cys Asp Cys Asp  
485 490 495

Gly Met Leu Asp Glu Glu Glu Phe Ala Leu Ala Lys His Leu Ile Lys  
500 505 510

Ile Lys Leu Asp Gly Tyr Glu Leu Pro Ser Ser Leu Pro Pro His Leu  
515 520 525

Val Pro Pro Ser His Arg Lys Ser Leu Pro Lys Ala Asp  
530 535 540

<210> 47  
<211> 1366  
<212> PRT  
<213> Homo sapiens

<400> 47

Met Leu Ala Val Gly Pro Ala Met Asp Arg Asp Tyr Pro Gln His Glu  
1 5 10 15

Pro Pro Pro Ala Gly Ser Leu Leu Tyr Ser Pro Pro Pro Leu Gln Ser  
20 25 30

Ala Met Leu His Cys Pro Tyr Trp Asn Thr Phe Ser Leu Pro Pro Tyr  
35 40 45

Pro Ala Phe Ser Ser Asp Ser Arg Pro Phe Met Ser Ser Ala Ser Phe  
50 55 60

Leu Gly Ser Gln Pro Cys Pro Asp Thr Ser Tyr Ala Pro Val Ala Thr  
65 70 75 80

Ala Ser Ser Leu Pro Pro Lys Thr Cys Asp Phe Ala Gln Asp Ser Ser  
85 90 95

Tyr Phe Glu Asp Phe Ser Asn Ile Ser Ile Phe Ser Ser Ser Val Asp  
100 105 110

Ser Leu Ser Asp Ile Val Asp Thr Pro Asp Phe Leu Pro Ala Asp Ser  
115 120 125

Leu Asn Gln Val Ser Thr Ile Trp Asp Asp Asn Pro Ala Pro Ser Thr  
130 135 140

His Asp Lys Leu Phe Gln Leu Ser Arg Pro Phe Ala Gly Phe Glu Asp  
145 150 155 160

Phe Leu Pro Ser His Ser Thr Pro Leu Leu Val Ser Tyr Gln Glu Gln  
165 170 175

Ser Val Gln Ser Gln Pro Glu Glu Glu Asp Glu Ala Glu Glu Glu  
180 185 190

Ala Glu Glu Leu Gly His Thr Glu Thr Tyr Ala Asp Tyr Val Pro Ser  
195 200 205

Lys Ser Lys Ile Gly Lys Gln His Pro Asp Arg Val Val Glu Thr Ser  
210 215 220

Thr Leu Ser Ser Val Pro Pro Pro Asp Ile Thr Tyr Thr Leu Ala Leu  
225 230 235 240

Pro Ser Asp Ser Gly Ala Leu Ser Ala Leu Gln Leu Glu Ala Ile Thr  
245 250 255

Tyr Ala Cys Gln Gln His Glu Val Leu Leu Pro Ser Gly Gln Arg Ala  
260 265 270

Gly Phe Leu Ile Gly Asp Gly Ala Gly Val Gly Lys Gly Arg Thr Val  
275 280 285

Ala Gly Val Ile Leu Glu Asn His Leu Arg Gly Arg Lys Lys Ala Leu  
290 295 300

Trp Phe Ser Val Ser Asn Asp Leu Lys Tyr Asp Ala Glu Arg Asp Leu  
305 310 315 320

Arg Asp Ile Glu Ala Thr Gly Ile Ala Val His Ala Leu Ser Lys Ile  
325 330 335

Lys Tyr Gly Asp Thr Thr Ser Glu Gly Val Leu Phe Ala Thr Tyr  
340 345 350

Ser Ala Leu Ile Gly Glu Ser Gln Ala Gly Gly Gln His Arg Thr Arg  
355 360 365

Leu Arg Gln Ile Leu Asp Trp Cys Gly Glu Ala Phe Glu Gly Val Ile  
370 375 380

Val Phe Asp Glu Cys His Lys Ala Lys Asn Ala Gly Ser Thr Lys Met  
385 390 395 400

Gly Lys Ala Val Leu Asp Leu Gln Asn Lys Leu Pro Leu Ala Arg Val  
405 410 415

Val Tyr Ala Ser Ala Thr Gly Ala Ser Glu Pro Arg Asn Met Ile Tyr

420

425

430

Met Ser Arg Leu Gly Ile Trp Gly Glu Gly Thr Pro Phe Arg Asn Phe  
 435 440 445

Glu Glu Phe Leu His Ala Ile Glu Lys Arg Gly Val Gly Ala Met Glu  
 450 455 460

Ile Val Ala Met Asp Met Lys Val Ser Gly Met Tyr Ile Ala Arg Gln  
 465 470 475 480

Leu Ser Phe Ser Gly Val Thr Phe Arg Ile Glu Glu Ile Pro Leu Ala  
 485 490 495

Pro Ala Phe Glu Cys Val Tyr Asn Arg Ala Ala Leu Leu Trp Ala Glu  
 500 505 510

Ala Leu Asn Val Phe Gln Gln Ala Ala Asp Trp Ile Gly Leu Glu Ser  
 515 520 525

Arg Lys Ser Leu Trp Gly Gln Phe Trp Ser Ala His Gln Arg Phe Phe  
 530 535 540

Lys Tyr Leu Cys Ile Ala Ala Lys Val Arg Arg Leu Val Glu Leu Ala  
 545 550 555 560

Arg Glu Glu Leu Ala Arg Asp Lys Cys Val Val Ile Gly Leu Gln Ser  
 565 570 575

Thr Gly Glu Ala Arg Thr Arg Glu Val Leu Gly Glu Asn Asp Gly His  
 580 585 590

Leu Asn Cys Phe Val Ser Ala Ala Glu Gly Val Phe Leu Ser Leu Ile  
 595 600 605

Gln Lys His Phe Pro Ser Thr Lys Arg Lys Arg Asp Arg Gly Ala Gly  
 610 615 620

Ser Lys Arg Lys Arg Arg Pro Arg Gly Arg Gly Ala Lys Ala Pro Arg  
 625 630 635 640

Leu Ala Cys Glu Thr Ala Gly Val Ile Arg Ile Ser Asp Asp Ser Ser  
 645 650 655

Thr Glu Ser Asp Pro Gly Leu Asp Ser Asp Phe Asn Ser Ser Pro Glu  
 660 665 670

Ser Leu Val Asp Asp Asp Val Val Ile Val Asp Ala Val Gly Leu Pro  
675 680 685

Ser Asp Asp Arg Gly Ser Leu Cys Leu Leu Gln Arg Asp Pro His Gly  
690 695 700

Pro Gly Val Leu Glu Arg Val Glu Arg Leu Lys Gln Asp Leu Leu Asp  
705 710 715 720

Lys Val Arg Arg Leu Gly Arg Glu Leu Pro Val Asn Thr Leu Asp Glu  
725 730 735

Leu Ile Asp Gln Leu Gly Gly Pro Gln Arg Val Ala Glu Met Thr Gly  
740 745 750

Arg Lys Gly Arg Val Val Ser Arg Pro Asp Gly Thr Val Ala Phe Glu  
755 760 765

Ser Arg Ala Glu Gln Gly Leu Ser Ile Asp His Val Asn Leu Arg Glu  
770 775 780

Lys Gln Arg Phe Met Ser Gly Glu Lys Leu Val Ala Ile Ile Ser Glu  
785 790 795 800

Ala Ser Ser Ser Gly Val Ser Leu Gln Ala Asp Arg Arg Val Gln Asn  
805 810 815

Gln Arg Arg Arg Val His Met Thr Leu Glu Leu Pro Trp Ser Ala Asp  
820 825 830

Arg Ala Ile Gln Gln Phe Gly Arg Thr His Arg Ser Asn Gln Val Ser  
835 840 845

Ala Pro Glu Tyr Val Phe Leu Ile Ser Glu Leu Ala Gly Glu Arg Arg  
850 855 860

Phe Ala Ser Ile Val Ala Lys Arg Leu Glu Ser Leu Gly Ala Leu Thr  
865 870 875 880

His Gly Asp Arg Arg Ala Thr Glu Ser Arg Asp Leu Ser Lys Tyr Asn  
885 890 895

Phe Glu Asn Lys Tyr Gly Thr Arg Ala Leu His Cys Val Leu Thr Thr  
900 905 910

Ile Leu Ser Gln Thr Glu Asn Lys Val Pro Val Pro Gln Gly Tyr Pro

915

920

925

Gly Gly Val Pro Thr Phe Phe Arg Asp Met Lys Gln Gly Leu Leu Ser  
 930 935 940

Val Gly Ile Gly Gly Arg Glu Ser Arg Asn Gly Cys Leu Asp Val Glu  
 945 950 955 960

Lys Asp Cys Ser Ile Thr Lys Phe Leu Asn Arg Ile Leu Gly Leu Glu  
 965 970 975

Val His Lys Gln Asn Ala Leu Phe Gln Tyr Phe Ser Asp Thr Phe Asp  
 980 985 990

His Leu Ile Glu Met Asp Lys Arg Glu Gly Lys Tyr Asp Met Gly Ile  
 995 1000 1005

Leu Asp Leu Ala Pro Gly Ile Glu Glu Ile Tyr Glu Glu Ser Gln  
 1010 1015 1020

Gln Val Phe Leu Ala Pro Gly His Pro Gln Asp Gly Gln Val Val  
 1025 1030 1035

Phe Tyr Lys Ile Ser Val Asp Arg Gly Leu Lys Trp Glu Asp Ala  
 1040 1045 1050

Phe Ala Lys Ser Leu Ala Leu Thr Gly Pro Tyr Asp Gly Phe Tyr  
 1055 1060 1065

Leu Ser Tyr Lys Val Arg Gly Asn Lys Pro Ser Cys Leu Leu Ala  
 1070 1075 1080

Glu Gln Asn Arg Gly Gln Phe Phe Thr Val Tyr Lys Pro Asn Ile  
 1085 1090 1095

Gly Arg Gln Ser Gln Leu Glu Ala Leu Asp Ser Leu Arg Arg Lys  
 1100 1105 1110

Phe His Arg Val Thr Ala Glu Glu Ala Lys Glu Pro Trp Glu Ser  
 1115 1120 1125

Gly Tyr Ala Leu Ser Leu Thr His Cys Ser His Ser Ala Trp Asn  
 1130 1135 1140

Arg His Cys Arg Leu Ala Gln Glu Gly Lys Asp Cys Leu Gln Gly  
 1145 1150 1155

Leu Arg Leu Arg His His Tyr Met Leu Cys Gly Ala Leu Leu Arg  
1160 1165 1170

Val Trp Gly Arg Ile Ala Ala Val Met Ala Asp Val Ser Ser Ser  
1175 1180 1185

Ser Tyr Leu Gln Ile Val Arg Leu Lys Thr Lys Asp Arg Lys Lys  
1190 1195 1200

Gln Val Gly Ile Lys Ile Pro Glu Gly Cys Val Arg Arg Val Leu  
1205 1210 1215

Gln Glu Leu Arg Leu Met Asp Ala Asp Val Lys Arg Arg Gln Ala  
1220 1225 1230

Pro Ala Leu Gly Cys Pro Ala Pro Pro Ala Pro Arg Pro Leu Ala  
1235 1240 1245

Leu Pro Cys Gly Pro Gly Glu Val Leu Asp Leu Thr Tyr Ser Pro  
1250 1255 1260

Pro Ala Glu Ala Phe Pro Pro Pro Pro His Phe Ser Phe Pro Ala  
1265 1270 1275

Pro Leu Ser Leu Asp Ala Gly Pro Gly Val Val Pro Leu Gly Thr  
1280 1285 1290

Pro Asp Ala Gln Ala Asp Pro Ala Ala Leu Ala His Gln Gly Cys  
1295 1300 1305

Asp Ile Asn Phe Lys Glu Val Leu Glu Asp Met Leu Arg Ser Leu  
1310 1315 1320

His Ala Gly Pro Pro Ser Glu Gly Ala Leu Gly Glu Gly Ala Gly  
1325 1330 1335

Ala Gly Gly Ala Ala Gly Gly Pro Glu Arg Gln Ser Val Ile  
1340 1345 1350

Gln Phe Ser Pro Pro Phe Pro Gly Ala Gln Ala Pro Leu  
1355 1360 1365

<210> 48

<211> .1392

<212> PRT

<213> Homo sapiens

<400> 48

Met Val Glu Pro Gly Gln Asp Leu Leu Leu Ala Ala Leu Ser Glu Ser  
1 5 10 15

Gly Ile Ser Pro Asn Asp Leu Phe Asp Ile Asp Gly Gly Asp Ala Gly  
20 25 30

Leu Ala Thr Pro Met Pro Thr Pro Ser Val Gln Gln Ser Val Pro Leu  
35 40 45

Ser Ala Leu Glu Leu Gly Leu Glu Thr Glu Ala Ala Val Pro Val Lys  
50 55 60

Gln Glu Pro Glu Thr Val Pro Thr Pro Ala Leu Leu Asn Val Arg Gln  
65 70 75 80

Pro Pro Ser Thr Thr Phe Val Leu Asn Gln Ile Asn His Leu Pro  
85 90 95

Pro Leu Gly Ser Thr Ile Val Met Thr Lys Thr Pro Pro Val Thr Thr  
100 105 110

Asn Arg Gln Thr Ile Thr Leu Thr Lys Phe Ile Gln Thr Thr Ala Ser  
115 120 125

Thr Arg Pro Ser Val Ser Ala Pro Thr Val Arg Asn Ala Met Thr Ser  
130 135 140

Ala Pro Ser Lys Asp Gln Val Gln Leu Lys Asp Leu Leu Lys Asn Asn  
145 150 155 160

Ser Leu Asn Glu Leu Met Lys Leu Lys Pro Pro Ala Asn Ile Ala Gln  
165 170 175

Pro Val Ala Thr Ala Ala Thr Asp Val Ser Asn Gly Thr Val Lys Lys  
180 185 190

Glu Ser Ser Asn Lys Glu Gly Ala Arg Met Trp Ile Asn Asp Met Lys  
195 200 205

Met Arg Ser Phe Ser Pro Thr Met Lys Val Pro Val Val Lys Glu Asp  
210 215 220

Asp Glu Pro Glu Glu Asp Glu Glu Met Gly His Ala Glu Thr  
225 230 235 240

Tyr Ala Glu Tyr Met Pro Ile Lys Leu Lys Ile Gly Leu Arg His Pro  
245 250 255

Asp Ala Val Val Glu Thr Ser Ser Leu Ser Ser Val Thr Pro Pro Asp  
260 265 270

Val Trp Tyr Lys Thr Ser Ile Ser Glu Glu Thr Ile Asp Asn Gly Trp  
275 280 285

Leu Ser Ala Leu Gln Leu Glu Ala Ile Thr Tyr Ala Ala Gln Gln His  
290 295 300

Glu Thr Phe Leu Pro Asn Gly Asp Arg Ala Gly Phe Leu Ile Gly Asp  
305 310 315 320

Gly Ala Gly Val Gly Lys Gly Arg Thr Ile Ala Gly Ile Ile Tyr Glu  
325 330 335

Asn Tyr Leu Leu Ser Arg Lys Arg Ala Leu Trp Phe Ser Val Ser Asn  
340 345 350

Asp Leu Lys Tyr Asp Ala Glu Arg Asp Leu Arg Asp Ile Gly Ala Lys  
355 360 365

Asn Ile Leu Val His Ser Leu Asn Lys Phe Lys Tyr Gly Lys Ile Ser  
370 375 380

Ser Lys His Asn Gly Ser Val Lys Lys Gly Val Ile Phe Ala Thr Tyr  
385 390 395 400

Ser Ser Leu Ile Gly Glu Ser Gln Ser Gly Gly Lys Tyr Lys Thr Arg  
405 410 415

Leu Lys Gln Leu Leu His Trp Cys Gly Asp Asp Phe Asp Gly Val Ile  
420 425 430

Val Phe Asp Glu Cys His Lys Ala Lys Asn Leu Cys Pro Val Gly Ser  
435 440 445

Ser Lys Pro Thr Lys Thr Gly Leu Ala Val Leu Glu Leu Gln Asn Lys  
450 455 460

Leu Pro Lys Ala Arg Val Val Tyr Ala Ser Ala Thr Gly Ala Ser Glu  
465 470 475 480

Pro Arg Asn Met Ala Tyr Met Asn Arg Leu Gly Ile Trp Gly Glu Gly  
485 490 495

Thr Pro Phe Arg Glu Phe Ser Asp Phe Ile Gln Ala Val Glu Arg Arg  
500 505 510

Gly Val Gly Ala Met Glu Ile Val Ala Met Asp Met Lys Leu Arg Gly  
515 520 525

Met Tyr Ile Ala Arg Gln Leu Ser Phe Thr Gly Val Thr Phe Lys Ile  
530 535 540

Glu Glu Val Leu Leu Ser Gln Ser Tyr Val Lys Met Tyr Asn Lys Ala  
545 550 555 560

Val Lys Leu Trp Val Ile Ala Arg Glu Arg Phe Gln Gln Ala Ala Asp  
565 570 575

Leu Ile Asp Ala Glu Gln Arg Met Lys Lys Ser Met Trp Gly Gln Phe  
580 585 590

Trp Ser Ala His Gln Arg Phe Phe Lys Tyr Leu Cys Ile Ala Ser Lys  
595 600 605

Val Lys Arg Val Val Gln Leu Ala Arg Glu Glu Ile Lys Asn Gly Lys  
610 615 620

Cys Val Val Ile Gly Leu Gln Ser Thr Gly Glu Ala Arg Thr Leu Glu  
625 630 635 640

Ala Leu Glu Glu Gly Gly Glu Leu Asn Asp Phe Val Ser Thr Ala  
645 650 655

Lys Gly Val Leu Gln Ser Leu Ile Glu Lys His Phe Pro Ala Pro Asp  
660 665 670

Arg Lys Lys Leu Tyr Ser Leu Leu Gly Ile Asp Leu Thr Ala Pro Ser  
675 680 685

Asn Asn Ser Ser Pro Arg Asp Ser Pro Cys Lys Glu Asn Lys Ile Lys  
690 695 700

Lys Arg Lys Gly Glu Glu Ile Thr Arg Glu Ala Lys Lys Ala Arg Lys  
705 710 715 720

Val Gly Gly Leu Thr Gly Ser Ser Ser Asp Asp Ser Gly Ser Glu Ser  
725 730 735

Asp Ala Ser Asp Asn Glu Glu Ser Asp Tyr Glu Ser Ser Lys Asn Met  
740 745 750

Ser Ser Gly Asp Asp Asp Phe Asn Pro Phe Leu Asp Glu Ser Asn  
755 760 765

Glu Asp Asp Glu Ser Asp Pro Trp Leu Ile Arg Lys Asp His Lys Lys  
770 775 780

Asn Lys Glu Lys Lys Lys Ser Ile Asp Pro Asp Ser Ile Gln  
785 790 795 800

Ser Ala Leu Leu Ala Ser Gly Leu Gly Ser Lys Arg Pro Ser Phe Ser  
805 810 815

Ser Thr Pro Val Ile Ser Pro Ala Pro Asn Ser Thr Pro Ala Asn Ser  
820 825 830

Asn Thr Asn Ser Asn Ser Ser Leu Ile Thr Ser Gln Asp Ala Val Glu  
835 840 845

Arg Ala Gln Gln Met Lys Lys Asp Leu Leu Asp Lys Leu Glu Lys Leu  
850 855 860

Ala Glu Asp Leu Pro Pro Asn Thr Leu Asp Glu Leu Ile Asp Glu Leu  
865 870 875 880

Gly Gly Pro Glu Asn Val Ala Glu Met Thr Gly Arg Lys Gly Arg Val  
885 890 895

Val Ser Asn Asp Asp Gly Ser Ile Ser Tyr Glu Ser Arg Ser Glu Leu  
900 905 910

Asp Val Pro Val Glu Ile Leu Asn Ile Thr Glu Lys Gln Arg Phe Met  
915 920 925

Asp Gly Asp Lys Asn Ile Ala Ile Ile Ser Glu Ala Ala Ser Ser Gly  
930 935 940

Ile Ser Leu Gln Ala Asp Arg Arg Ala Lys Asn Gln Arg Arg Arg Val  
945 950 955 960

His Met Thr Leu Glu Leu Pro Trp Ser Ala Asp Arg Ala Ile Gln Gln  
965 970 975

Phe Gly Arg Thr His Arg Ser Asn Gln Val Thr Ala Pro Glu Tyr Val  
980 985 990

Phe Leu Ile Ser Glu Leu Ala Gly Glu Gln Arg Phe Ala Ser Ile Val  
995 1000 1005

Ala Lys Arg Leu Glu Ser Leu Gly Ala Leu Thr His Gly Asp Arg  
1010 1015 1020

Arg Ala Thr Glu Ser Arg Asp Leu Ser Arg Phe Asn Phe Asp Asn  
1025 1030 1035

Lys Tyr Gly Arg Asn Ala Leu Glu Ile Val Met Lys Ser Ile Val  
1040 1045 1050

Asn Leu Asp Ser Pro Met Val Ser Pro Pro Pro Asp Tyr Pro Gly  
1055 1060 1065

Glu Phe Phe Lys Asp Val Arg Gln Gly Leu Ile Gly Val Gly Leu  
1070 1075 1080

Ile Asn Val Glu Asp Arg Ser Gly Ile Leu Thr Leu Asp Lys Asp  
1085 1090 1095

Tyr Asn Asn Ile Gly Lys Phe Leu Asn Arg Ile Leu Gly Met Glu  
1100 1105 1110

Val His Gln Gln Asn Ala Leu Phe Gln Tyr Phe Ala Asp Thr Leu  
1115 1120 1125

Thr Ala Val Val Gln Asn Ala Lys Lys Asn Gly Arg Tyr Asp Met  
1130 1135 1140

Gly Ile Leu Asp Leu Gly Ser Gly Asp Glu Lys Val Arg Lys Ser  
1145 1150 1155

Asp Val Lys Lys Phe Leu Thr Pro Gly Tyr Ser Thr Ser Gly His  
1160 1165 1170

Val Glu Leu Tyr Thr Ile Ser Val Glu Arg Gly Met Ser Trp Glu  
1175 1180 1185

Glu Ala Thr Lys Ile Trp Ala Glu Leu Thr Gly Pro Asp Asp Gly  
1190 1195 1200

Phe Tyr Leu Ser Leu Gln Ile Arg Asn Asn Lys Lys Thr Ala Ile  
1205 1210 1215

Leu Val Lys Glu Val Asn Pro Lys Lys Lys Leu Phe Leu Val Tyr  
1220 1225 1230

Arg Pro Asn Thr Gly Lys Gln Leu Lys Leu Glu Ile Tyr Ala Asp  
1235 1240 1245

Leu Lys Lys Lys Tyr Lys Lys Val Val Ser Asp Asp Ala Leu Met  
1250 1255 1260

His Trp Leu Asp Gln Tyr Asn Ser Ser Ala Asp Thr Cys Thr His  
1265 1270 1275

Ala Tyr Trp Arg Gly Asn Cys Lys Lys Ala Ser Leu Gly Leu Val  
1280 1285 1290

Cys Glu Ile Gly Leu Arg Cys Arg Thr Tyr Tyr Val Leu Cys Gly  
1295 1300 1305

Ser Val Leu Ser Val Trp Thr Lys Val Glu Gly Val Leu Ala Ser  
1310 1315 1320

Val Ser Gly Thr Asn Val Lys Met Gln Ile Val Arg Leu Arg Thr  
1325 1330 1335

Glu Asp Gly Gln Arg Ile Val Gly Leu Ile Ile Pro Ala Asn Cys  
1340 1345 1350

Val Ser Pro Leu Val Asn Leu Leu Ser Thr Ser Asp Gln Ser Gln  
1355 1360 1365

Gln Leu Ala Val Gln Gln Lys Gln Leu Trp Gln Gln His His Pro  
1370 1375 1380

Gln Ser Ile Thr Asn Leu Ser Asn Ala  
1385 1390

<210> 49  
<211> 1327  
<212> PRT  
<213> Homo sapiens

<400> 49

Met Ala Ala Ser Arg Arg Ser Gln His His His His His Gln Gln  
1 5 10 15

Gln Leu Gln Pro Ala Pro Gly Ala Ser Ala Pro Pro Pro Pro Pro  
20 25 30

Pro Pro Leu Ser Pro Gly Leu Ala Pro Gly Thr Thr Pro Ala Ser Pro  
35 40 45

Thr Ala Ser Gly Leu Ala Pro Phe Ala Ser Pro Arg His Gly Leu Ala  
50 55 60

Leu Pro Glu Gly Asp Gly Ser Arg Asp Pro Pro Asp Arg Pro Arg Ser  
65 70 75 80

Pro Asp Pro Val Asp Gly Thr Ser Cys Cys Ser Thr Thr Ser Thr Ile  
85 90 95

Cys Thr Val Ala Ala Ala Pro Val Val Pro Ala Val Ser Thr Ser Ser  
100 105 110

Ala Ala Gly Val Ala Pro Asn Pro Ala Gly Ser Gly Ser Asn Asn Ser  
115 120 125

Pro Ser Ser Ser Ser Ser Pro Thr Ser Ser Ser Ser Ser Pro Ser  
130 135 140

Ser Pro Gly Ser Ser Leu Ala Glu Ser Pro Glu Ala Ala Gly Val Ser  
145 150 155 160

Ser Thr Ala Pro Leu Gly Pro Gly Ala Ala Gly Pro Gly Thr Gly Val  
165 170 175

Pro Ala Val Ser Gly Ala Leu Arg Glu Leu Leu Glu Ala Cys Arg Asn  
180 185 190

Gly Asp Val Ser Arg Val Lys Arg Leu Val Asp Ala Ala Asn Val Asn  
195 200 205

Ala Lys Asp Met Ala Gly Arg Lys Ser Ser Pro Leu His Phe Ala Ala  
210 215 220

Gly Phe Gly Arg Lys Asp Val Val Glu His Leu Leu Gln Met Gly Ala  
225 230 235 240

Asn Val His Ala Arg Asp Asp Gly Gly Leu Ile Pro Leu His Asn Ala  
245 250 255

Cys Ser Phe Gly His Ala Glu Val Val Ser Leu Leu Leu Cys Gln Gly  
260 265 270

Ala Asp Pro Asn Ala Arg Asp Asn Trp Asn Tyr Thr Pro Leu His Glu

275

280

285

Ala Ala Ile Lys Gly Lys Ile Asp Val Cys Ile Val Leu Leu Gln His  
 290 295 300

Gly Ala Asp Pro Asn Ile Arg Asn Thr Asp Gly Lys Ser Ala Leu Asp  
 305 310 315 320

Leu Ala Asp Pro Ser Ala Lys Ala Val Leu Thr Gly Glu Tyr Lys Lys  
 325 330 335

Asp Glu Leu Leu Glu Ala Ala Arg Ser Gly Asn Glu Glu Lys Leu Met  
 340 345 350

Ala Leu Leu Thr Pro Leu Asn Val Asn Cys His Ala Ser Asp Gly Arg  
 355 360 365

Lys Ser Thr Pro Leu His Leu Ala Ala Gly Tyr Asn Arg Val Arg Ile  
 370 375 380

Val Gln Leu Leu Leu Gln His Gly Ala Asp Val His Ala Lys Asp Lys  
 385 390 395 400

Gly Gly Leu Val Pro Leu His Asn Ala Cys Ser Tyr Gly His Tyr Glu  
 405 410 415

Val Thr Glu Leu Leu Leu Lys His Gly Ala Cys Val Asn Ala Met Asp  
 420 425 430

Leu Trp Gln Phe Thr Pro Leu His Glu Ala Ala Ser Lys Asn Arg Val  
 435 440 445

Glu Val Cys Ser Leu Leu Leu Ser His Gly Ala Asp Pro Thr Leu Val  
 450 455 460

Asn Cys His Gly Lys Ser Ala Val Asp Met Ala Pro Thr Pro Glu Leu  
 465 470 475 480

Arg Glu Arg Leu Thr Tyr Glu Phe Lys Gly His Ser Leu Leu Gln Ala  
 485 490 495

Ala Arg Glu Ala Asp Leu Ala Lys Val Lys Lys Thr Leu Ala Leu Glu  
 500 505 510

Ile Ile Asn Phe Lys Gln Pro Gln Ser His Glu Thr Ala Leu His Cys  
 515 520 525

Ala Val Ala Ser Leu His Pro Lys Arg Lys Gln Val Thr Glu Leu Leu  
530 535 540

Leu Arg Lys Gly Ala Asn Val Asn Glu Lys Asn Lys Asp Phe Met Thr  
545 550 555 560

Pro Leu His Val Ala Ala Glu Arg Ala His Asn Asp Val Met Glu Val  
565 570 575

Leu His Lys His Gly Ala Lys Met Asn Ala Leu Asp Thr Leu Gly Gln  
580 585 590

Thr Ala Leu His Arg Ala Ala Leu Ala Gly His Leu Gln Thr Cys Arg  
595 600 605

Leu Leu Leu Ser Tyr Gly Ser Asp Pro Ser Ile Ile Ser Leu Gln Gly  
610 615 620

Phe Thr Ala Ala Gln Met Gly Asn Glu Ala Val Gln Gln Ile Leu Ser  
625 630 635 640

Glu Ser Thr Pro Ile Arg Thr Ser Asp Val Asp Tyr Arg Leu Leu Glu  
645 650 655

Ala Ser Lys Ala Gly Asp Leu Glu Thr Val Lys Gln Leu Cys Ser Ser  
660 665 670

Gln Asn Val Asn Cys Arg Asp Leu Glu Gly Arg His Ser Thr Pro Leu  
675 680 685

His Phe Ala Ala Gly Tyr Asn Arg Val Ser Val Val Glu Tyr Leu Leu  
690 695 700

His His Gly Ala Asp Val His Ala Lys Asp Lys Gly Gly Leu Val Pro  
705 710 715 720

Leu His Asn Ala Cys Ser Tyr Gly His Tyr Glu Val Ala Glu Leu Leu  
725 730 735

Val Arg His Gly Ala Ser Val Asn Val Ala Asp Leu Trp Lys Phe Thr  
740 745 750

Pro Leu His Glu Ala Ala Ala Lys Gly Lys Tyr Glu Ile Cys Lys Leu  
755 760 765

Leu Leu Lys His Gly Ala Asp Pro Thr Lys Lys Asn Arg Asp Gly Asn

770

775

780

Thr Pro Leu Asp Leu Val Lys Glu Gly Asp Thr Asp Ile Gln Asp Leu  
 785 790 795 800

Leu Lys Gly Asp Ala Ala Leu Leu Asp Ala Ala Lys Lys Gly Cys Leu  
 805 810 815

Ala Arg Val Gln Lys Leu Cys Thr Pro Glu Asn Ile Asn Cys Arg Asp  
 820 825 830

Thr Gln Gly Arg Asn Ser Thr Pro Leu His Leu Ala Ala Gly Tyr Asn  
 835 840 845

Asn Leu Glu Val Ala Glu Tyr Leu Leu Glu His Gly Ala Asp Val Asn  
 850 855 860

Ala Gln Asp Lys Gly Gly Leu Ile Pro Leu His Asn Ala Ala Ser Tyr  
 865 870 875 880

Gly His Val Asp Ile Ala Ala Leu Leu Ile Lys Tyr Asn Thr Cys Val  
 885 890 895

Asn Ala Thr Asp Lys Trp Ala Phe Thr Pro Leu His Glu Ala Ala Gln  
 900 905 910

Lys Gly Arg Thr Gln Leu Cys Ala Leu Leu Leu Ala His Gly Ala Asp  
 915 920 925

Pro Thr Met Lys Asn Gln Glu Gly Gln Thr Pro Leu Asp Leu Ala Thr  
 930 935 940

Ala Asp Asp Ile Arg Ala Leu Leu Ile Asp Ala Met Pro Pro Glu Ala  
 945 950 955 960

Leu Pro Thr Cys Phe Lys Pro Gln Ala Thr Val Val Ser Ala Ser Leu  
 965 970 975

Ile Ser Pro Ala Ser Thr Pro Ser Cys Leu Ser Ala Ala Ser Ser Ile  
 980 985 990

Asp Asn Leu Thr Gly Pro Leu Ala Glu Leu Ala Val Gly Gly Ala Ser  
 995 1000 1005

Asn Ala Gly Asp Gly Ala Ala Gly Thr Glu Arg Lys Glu Gly Glu  
 1010 1015 1020

Val Ala Gly Leu Asp Met Asn Ile Ser Gln Phe Leu Lys Ser Leu  
1025 1030 1035

Gly Leu Glu His Leu Arg Asp Ile Phe Glu Thr Glu Gln Ile Thr  
1040 1045 1050

Leu Asp Val Leu Ala Asp Met Gly His Glu Glu Leu Lys Glu Ile  
1055 1060 1065

Gly Ile Asn Ala Tyr Gly His Arg His Lys Leu Ile Lys Gly Val  
1070 1075 1080

Glu Arg Leu Leu Gly Gly Gln Gln Gly Thr Asn Pro Tyr Leu Thr  
1085 1090 1095

Phe His Cys Val Asn Gln Gly Thr Ile Leu Leu Asp Leu Ala Pro  
1100 1105 1110

Glu Asp Lys Glu Tyr Gln Ser Val Glu Glu Glu Met Gln Ser Thr  
1115 1120 1125

Ile Arg Glu His Arg Asp Gly Gly Asn Ala Gly Gly Ile Phe Asn  
1130 1135 1140

Arg Tyr Asn Val Ile Arg Ile Gln Lys Val Val Asn Lys Lys Leu  
1145 1150 1155

Arg Glu Arg Phe Cys His Arg Gln Lys Glu Val Ser Glu Glu Asn  
1160 1165 1170

His Asn His His Asn Glu Arg Met Leu Phe His Gly Ser Pro Phe  
1175 1180 1185

Ile Asn Ala Ile Ile His Lys Gly Phe Asp Glu Arg His Ala Tyr  
1190 1195 1200

Ile Gly Gly Met Phe Gly Ala Gly Ile Tyr Phe Ala Glu Asn Ser  
1205 1210 1215

Ser Lys Ser Asn Gln Tyr Val Tyr Gly Ile Gly Gly Gly Thr Gly  
1220 1225 1230

Cys Pro Thr His Lys Asp Arg Ser Cys Tyr Ile Cys His Arg Gln  
1235 1240 1245

Met Leu Phe Cys Arg Val Thr Leu Gly Lys Ser Phe Leu Gln Phe

1250

1255

1260

Ser Thr Met Lys Met Ala His Ala Pro Pro Gly His His Ser Val  
 1265 1270 1275

Ile Gly Arg Pro Ser Val Asn Gly Leu Ala Tyr Ala Glu Tyr Val  
 1280 1285 1290

Ile Tyr Arg Gly Glu Gln Ala Tyr Pro Glu Tyr Leu Ile Thr Tyr  
 1295 1300 1305

Gln Ile Met Lys Pro Glu Ala Pro Ser Gln Thr Ala Thr Ala Ala  
 1310 1315 1320

Glu Gln Lys Thr  
 1325

<210> 50  
 <211> 1166  
 <212> PRT  
 <213> Homo sapiens

<400> 50

Met Ser Gly Arg Arg Cys Ala Gly Gly Ala Ala Cys Ala Ser Ala  
 1 5 10 15

Ala Ala Glu Ala Val Glu Pro Ala Ala Arg Glu Leu Phe Glu Ala Cys  
 20 25 30

Arg Asn Gly Asp Val Glu Arg Val Lys Arg Leu Val Thr Pro Glu Lys  
 35 40 45

Val Asn Ser Arg Asp Thr Ala Gly Arg Lys Ser Thr Pro Leu His Phe  
 50 55 60

Ala Ala Gly Phe Gly Arg Lys Asp Val Val Glu Tyr Leu Leu Gln Asn  
 65 70 75 80

Gly Ala Asn Val Gln Ala Arg Asp Asp Gly Gly Leu Ile Pro Leu His  
 85 90 95

Asn Ala Cys Ser Phe Gly His Ala Glu Val Val Asn Leu Leu Arg  
 100 105 110

His Gly Ala Asp Pro Asn Ala Arg Asp Asn Trp Asn Tyr Thr Pro Leu  
 115 120 125

His Glu Ala Ala Ile Lys Gly Lys Ile Asp Val Cys Ile Val Leu Leu  
130 135 140

Gln His Gly Ala Glu Pro Thr Ile Arg Asn Thr Asp Gly Arg Thr Ala  
145 150 155 160

Leu Asp Leu Ala Asp Pro Ser Ala Lys Ala Val Leu Thr Gly Glu Tyr  
165 170 175

Lys Lys Asp Glu Leu Leu Glu Ser Ala Arg Ser Gly Asn Glu Glu Lys  
180 185 190

Met Met Ala Leu Leu Thr Pro Leu Asn Val Asn Cys His Ala Ser Asp  
195 200 205

Gly Arg Lys Ser Thr Pro Leu His Leu Ala Ala Gly Tyr Asn Arg Val  
210 215 220

Lys Ile Val Gln Leu Leu Leu Gln His Gly Ala Asp Val His Ala Lys  
225 230 235 240

Asp Lys Gly Asp Leu Val Pro Leu His Asn Ala Cys Ser Tyr Gly His  
245 250 255

Tyr Glu Val Thr Glu Leu Leu Val Lys His Gly Ala Cys Val Asn Ala  
260 265 270

Met Asp Leu Trp Gln Phe Thr Pro Leu His Glu Ala Ala Ser Lys Asn  
275 280 285

Arg Val Glu Val Cys Ser Leu Leu Leu Ser Tyr Gly Ala Asp Pro Thr  
290 295 300

Leu Leu Asn Cys His Asn Lys Ser Ala Ile Asp Leu Ala Pro Thr Pro  
305 310 315 320

Gln Leu Lys Glu Arg Leu Ala Tyr Glu Phe Lys Gly His Ser Leu Leu  
325 330 335

Gln Ala Ala Arg Glu Ala Asp Val Thr Arg Ile Lys Lys His Leu Ser  
340 345 350

Leu Glu Met Val Asn Phe Lys His Pro Gln Thr His Glu Thr Ala Leu  
355 360 365

His Cys Ala Ala Ala Ser Pro Tyr Pro Lys Arg Lys Gln Ile Cys Glu  
370 375 380

Leu Leu Leu Arg Lys Gly Ala Asn Ile Asn Glu Lys Thr Lys Glu Phe  
385 390 395 400

Leu Thr Pro Leu His Val Ala Ser Glu Lys Ala His Asn Asp Val Val  
405 410 415

Glu Val Val Val Lys His Glu Ala Lys Val Asn Ala Leu Asp Asn Leu  
420 425 430

Gly Gln Thr Ser Leu His Arg Ala Ala Tyr Cys Gly His Leu Gln Thr  
435 440 445

Cys Arg Leu Leu Leu Ser Tyr Gly Cys Asp Pro Asn Ile Ile Ser Leu  
450 455 460

Gln Gly Phe Thr Ala Leu Gln Met Gly Asn Glu Asn Val Gln Gln Leu  
465 470 475 480

Leu Gln Glu Gly Ile Ser Leu Gly Asn Ser Glu Ala Asp Arg Gln Leu  
485 490 495

Leu Glu Ala Ala Lys Ala Gly Asp Val Glu Thr Val Lys Lys Leu Cys  
500 505 510

Thr Val Gln Ser Val Asn Cys Arg Asp Ile Glu Gly Arg Gln Ser Thr  
515 520 525

Pro Leu His Phe Ala Ala Gly Tyr Asn Arg Val Ser Val Val Glu Tyr  
530 535 540

Leu Leu Gln His Gly Ala Asp Val His Ala Lys Asp Lys Gly Gly Leu  
545 550 555 560

Val Pro Leu His Asn Ala Cys Ser Tyr Gly His Tyr Glu Val Ala Glu  
565 570 575

Leu Leu Val Lys His Gly Ala Val Val Asn Val Ala Asp Leu Trp Lys  
580 585 590

Phe Thr Pro Leu His Glu Ala Ala Ala Lys Gly Lys Tyr Glu Ile Cys  
595 600 605

Lys Leu Leu Leu Gln His Gly Ala Asp Pro Thr Lys Lys Asn Arg Asp  
610 615 620

Gly Asn Thr Pro Leu Asp Leu Val Lys Asp Gly Asp Thr Asp Ile Gln  
625 630 635 640

Asp Leu Leu Arg Gly Asp Ala Ala Leu Leu Asp Ala Ala Lys Lys Gly  
645 650 655

Cys Leu Ala Arg Val Lys Lys Leu Ser Ser Pro Asp Asn Val Asn Cys  
660 665 670

Arg Asp Thr Gln Gly Arg His Ser Thr Pro Leu His Leu Ala Ala Gly  
675 680 685

Tyr Asn Asn Leu Glu Val Ala Glu Tyr Leu Leu Gln His Gly Ala Asp  
690 695 700

Val Asn Ala Gln Asp Lys Gly Gly Leu Ile Pro Leu His Asn Ala Ala  
705 710 715 720

Ser Tyr Gly His Val Asp Val Ala Ala Leu Leu Ile Lys Tyr Asn Ala  
725 730 735

Cys Val Asn Ala Thr Asp Lys Trp Ala Phe Thr Pro Leu His Glu Ala  
740 745 750

Ala Gln Lys Gly Arg Thr Gln Leu Cys Ala Leu Leu Leu Ala His Gly  
755 760 765

Ala Asp Pro Thr Leu Lys Asn Gln Glu Gly Gln Thr Pro Leu Asp Leu  
770 775 780

Val Ser Ala Asp Asp Val Ser Ala Leu Leu Thr Ala Ala Met Pro Pro  
785 790 795 800

Ser Ala Leu Pro Ser Cys Tyr Lys Pro Gln Val Leu Asn Gly Val Arg  
805 810 815

Ser Pro Gly Ala Thr Ala Asp Ala Leu Ser Ser Gly Pro Ser Ser Pro  
820 825 830

Ser Ser Leu Ser Ala Ala Ser Ser Leu Asp Asn Leu Ser Gly Ser Phe  
835 840 845

Ser Glu Leu Ser Ser Val Val Ser Ser Ser Gly Thr Glu Gly Ala Ser  
850 855 860

Ser Leu Glu Lys Lys Glu Val Pro Gly Val Asp Phe Ser Ile Thr Gln  
865 870 875 880

Phe Val Arg Asn Leu Gly Leu Glu His Leu Met Asp Ile Phe Glu Arg  
885 890 895

Glu Gln Ile Thr Leu Asp Val Leu Val Glu Met Gly His Lys Glu Leu  
900 905 910

Lys Glu Ile Gly Ile Asn Ala Tyr Gly His Arg His Lys Leu Ile Lys  
915 920 925

Gly Val Glu Arg Leu Ile Ser Gly Gln Gln Gly Leu Asn Pro Tyr Leu  
930 935 940

Thr Leu Asn Thr Ser Gly Ser Gly Thr Ile Leu Ile Asp Leu Ser Pro  
945 950 955 960

Asp Asp Lys Glu Phe Gln Ser Val Glu Glu Glu Met Gln Ser Thr Val  
965 970 975

Arg Glu His Arg Asp Gly Gly His Ala Gly Gly Ile Phe Asn Arg Tyr  
980 985 990

Asn Ile Leu Lys Ile Gln Lys Val Cys Asn Lys Lys Leu Trp Glu Arg  
995 1000 1005

Tyr Thr His Arg Arg Lys Glu Val Ser Glu Glu Asn His Asn His  
1010 1015 1020

Ala Asn Glu Arg Met Leu Phe His Gly Ser Pro Phe Val Asn Ala  
1025 1030 1035

Ile Ile His Lys Gly Phe Asp Glu Arg His Ala Tyr Ile Gly Gly  
1040 1045 1050

Met Phe Gly Ala Gly Ile Tyr Phe Ala Glu Asn Ser Ser Lys Ser  
1055 1060 1065

Asn Gln Tyr Val Tyr Gly Ile Gly Gly Gly Thr Gly Cys Pro Val  
1070 1075 1080

His Lys Asp Arg Ser Cys Tyr Ile Cys His Arg Gln Leu Leu Phe  
1085 1090 1095

Cys Arg Val Thr Leu Gly Lys Ser Phe Leu Gln Phe Ser Ala Met  
1100 1105 1110

Lys Met Ala His Ser Pro Pro Gly His His Ser Val Thr Gly Arg  
1115 1120 1125

Pro Ser Val Asn Gly Leu Ala Leu Ala Glu Tyr Val Ile Tyr Arg  
1130 1135 1140

Gly Glu Gln Ala Tyr Pro Glu Tyr Leu Ile Thr Tyr Gln Ile Met  
1145 1150 1155

Arg Pro Glu Gly Met Val Asp Gly  
1160 1165

<210> 51

<211> 1243

<212> PRT

<213> Homo sapiens

<400> 51

Met Ser Glu Ala Pro Arg Phe Phe Val Gly Pro Glu Asp Thr Glu Ile  
1 5 10 15

Asn Pro Gly Asn Tyr Arg His Phe Phe His His Ala Asp Glu Asp Asp  
20 25 30

Glu Glu Glu Asp Asp Ser Pro Pro Glu Arg Gln Ile Val Val Gly Ile  
35 40 45

Cys Ser Met Ala Lys Lys Ser Lys Ser Lys Pro Met Lys Glu Ile Leu  
50 55 60

Glu Arg Ile Ser Leu Phe Lys Tyr Ile Thr Val Val Val Phe Glu Glu  
65 70 75 80

Glu Val Ile Leu Asn Glu Pro Val Glu Asn Trp Pro Leu Cys Asp Cys  
85 90 95

Leu Ile Ser Phe His Ser Lys Gly Phe Pro Leu Asp Lys Ala Val Ala  
100 105 110

Tyr Ala Lys Leu Arg Asn Pro Phe Val Ile Asn Asp Leu Asn Met Gln  
115 120 125

Tyr Leu Ile Gln Asp Arg Arg Glu Val Tyr Ser Ile Leu Gln Ala Glu  
130 135 140

Gly Ile Leu Leu Pro Arg Tyr Ala Ile Leu Asn Arg Asp Pro Asn Asn  
145 150 155 160

Pro Lys Glu Cys Asn Leu Ile Glu Gly Glu Asp His Val Glu Val Asn  
165 170 175

Gly Glu Val Phe Gln Lys Pro Phe Val Glu Lys Pro Val Ser Ala Glu  
180 185 190

Asp His Asn Val Tyr Ile Tyr Tyr Pro Thr Ser Ala Gly Gly Ser  
195 200 205

Gln Arg Leu Phe Arg Lys Ile Gly Ser Arg Ser Ser Val Tyr Ser Pro  
210 215 220

Glu Ser Asn Val Arg Lys Thr Gly Ser Tyr Ile Tyr Glu Glu Phe Met  
225 230 235 240

Pro Thr Asp Gly Thr Asp Val Lys Val Tyr Thr Val Gly Pro Asp Tyr  
245 250 255

Ala His Ala Glu Ala Arg Lys Ser Pro Ala Leu Asp Gly Lys Val Glu  
260 265 270

Arg Asp Ser Glu Gly Lys Glu Val Arg Tyr Pro Val Ile Leu Asn Ala  
275 280 285

Arg Glu Lys Leu Ile Ala Trp Lys Val Cys Leu Ala Phe Lys Gln Thr  
290 295 300

Val Cys Gly Phe Asp Leu Leu Arg Ala Asn Gly Gln Ser Tyr Val Cys  
305 310 315 320

Asp Val Asn Gly Phe Ser Phe Val Lys Asn Ser Met Lys Tyr Tyr Asp  
325 330 335

Asp Cys Ala Lys Ile Leu Gly Asn Ile Val Met Arg Glu Leu Ala Pro  
340 345 350

Gln Phe His Ile Pro Trp Ser Ile Pro Leu Glu Ala Glu Asp Ile Pro  
355 360 365

Ile Val Pro Thr Thr Ser Gly Thr Met Met Glu Leu Arg Cys Val Ile  
370 375 380

Ala Val Ile Arg His Gly Asp Arg Thr Pro Lys Gln Lys Met Lys Met  
385 390 395 400

Glu Val Arg His Gln Lys Phe Phe Asp Leu Phe Glu Lys Cys Asp Gly

405

410

415

Tyr Lys Ser Gly Lys Leu Lys Leu Lys Pro Lys Gln Leu Gln Glu  
 420 425 430

Val Leu Asp Ile Ala Arg Gln Leu Leu Met Glu Leu Gly Gln Asn Asn  
 435 440 445

Asp Ser Glu Ile Glu Glu Asn Lys Pro Lys Leu Glu Gln Leu Lys Thr  
 450 455 460

Val Leu Glu Met Tyr Gly His Phe Ser Gly Ile Asn Arg Lys Val Gln  
 465 470 475 480

Leu Thr Tyr Leu Pro His Gly Cys Pro Lys Thr Ser Ser Glu Glu Glu  
 485 490 495

Asp Ser Arg Arg Glu Glu Pro Ser Leu Leu Leu Val Leu Lys Trp Gly  
 500 505 510

Gly Glu Leu Thr Pro Ala Gly Arg Val Gln Ala Glu Glu Leu Gly Arg  
 515 520 525

Ala Phe Arg Cys Met Tyr Pro Gly Gly Gln Gly Asp Tyr Ala Gly Phe  
 530 535 540

Pro Gly Cys Gly Leu Leu Arg Leu His Ser Thr Tyr Arg His Asp Leu  
 545 550 555 560

Lys Ile Tyr Ala Ser Asp Glu Gly Arg Val Gln Met Thr Ala Ala Ala  
 565 570 575

Phe Ala Lys Gly Leu Leu Ala Leu Glu Gly Glu Leu Thr Pro Ile Leu  
 580 585 590

Val Gln Met Val Lys Ser Ala Asn Met Asn Gly Leu Leu Asp Ser Asp  
 595 600 605

Ser Asp Ser Leu Ser Ser Cys Gln Gln Arg Val Lys Ala Arg Leu His  
 610 615 620

Glu Ile Leu Gln Lys Asp Arg Asp Phe Thr Ala Glu Asp Tyr Glu Lys  
 625 630 635 640

Leu Thr Pro Ser Gly Ser Ile Ser Leu Ile Lys Ser Met His Leu Ile  
 645 650 655

Lys Asn Pro Val Lys Thr Cys Asp Lys Val Tyr Ser Leu Ile Gln Ser  
660 665 670

Leu Thr Ser Gln Ile Arg His Arg Met Glu Asp Pro Lys Ser Ser Asp  
675 680 685

Ile Gln Leu Tyr His Ser Glu Thr Leu Glu Leu Met Leu Arg Arg Trp  
690 695 700

Ser Lys Leu Glu Lys Asp Phe Lys Thr Lys Asn Gly Arg Tyr Asp Ile  
705 710 715 720

Ser Lys Ile Pro Asp Ile Tyr Asp Cys Ile Lys Tyr Asp Val Gln His  
725 730 735

Asn Gly Ser Leu Lys Leu Glu Asn Thr Met Glu Leu Tyr Arg Leu Ser  
740 745 750

Lys Ala Leu Ala Asp Ile Val Ile Pro Gln Glu Tyr Gly Ile Thr Lys  
755 760 765

Ala Glu Lys Leu Glu Ile Ala Lys Gly Tyr Cys Thr Pro Leu Val Arg  
770 775 780

Lys Ile Arg Ser Asp Leu Gln Arg Thr Gln Asp Asp Asp Thr Val Asn  
785 790 795 800

Lys Leu His Pro Val Tyr Ser Arg Gly Val Leu Ser Pro Glu Arg His  
805 810 815

Val Arg Thr Arg Leu Tyr Phe Thr Ser Glu Ser His Val His Ser Leu  
820 825 830

Leu Ser Ile Leu Arg Tyr Gly Ala Leu Cys Asn Glu Ser Lys Asp Glu  
835 840 845

Gln Trp Lys Arg Ala Met Asp Tyr Leu Asn Val Val Asn Glu Leu Asn  
850 855 860

Tyr Met Thr Gln Ile Val Ile Met Leu Tyr Glu Asp Pro Asn Lys Asp  
865 870 875 880

Leu Ser Ser Glu Glu Arg Phe His Val Glu Leu His Phe Ser Pro Gly  
885 890 895

Ala Lys Gly Cys Glu Glu Asp Lys Asn Leu Pro Ser Gly Tyr Gly Tyr

900

905

910

Arg Pro Ala Ser Arg Glu Asn Glu Gly Arg Arg Pro Phe Lys Ile Asp  
915 920 925

Asn Asp Asp Glu Pro His Thr Ser Lys Arg Asp Glu Val Asp Arg Ala  
930 935 940

Val Ile Leu Phe Lys Pro Met Val Ser Glu Pro Ile His Ile His Arg  
945 950 955 960

Lys Ser Pro Leu Pro Arg Ser Arg Lys Thr Ala Thr Asn Asp Glu Glu  
965 970 975

Ser Pro Leu Ser Val Ser Ser Pro Glu Gly Thr Gly Thr Trp Leu His  
980 985 990

Tyr Thr Ser Gly Val Gly Thr Gly Arg Arg Arg Arg Arg Ser Gly Glu  
995 1000 1005

Gln Ile Thr Ser Ser Pro Val Ser Pro Lys Ser Leu Ala Phe Thr  
1010 1015 1020

Ser Ser Ile Phe Gly Ser Trp Gln Gln Val Val Ser Glu Asn Ala  
1025 1030 1035

Asn Tyr Leu Arg Thr Pro Arg Thr Leu Val Glu Gln Lys Gln Asn  
1040 1045 1050

Pro Thr Val Gly Ser His Cys Ala Gly Leu Phe Ser Thr Ser Val  
1055 1060 1065

Leu Gly Gly Ser Ser Ser Ala Pro Asn Leu Gln Asp Tyr Ala Arg  
1070 1075 1080

Thr His Arg Lys Lys Leu Thr Ser Ser Gly Cys Ile Asp Asp Ala  
1085 1090 1095

Thr Arg Gly Ser Ala Val Lys Arg Phe Ser Ile Ser Phe Ala Arg  
1100 1105 1110

His Pro Thr Asn Gly Phe Glu Leu Tyr Ser Met Val Pro Ser Ile  
1115 1120 1125

Cys Pro Leu Glu Thr Leu His Asn Ala Leu Ser Leu Lys Gln Val  
1130 1135 1140

Asp Glu Phe Leu Ala Ser Ile Ala Ser Pro Ser Ser Asp Val Pro  
1145 1150 1155

Arg Lys Thr Ala Glu Ile Ser Ser Thr Ala Leu Arg Ser Ser Pro  
1160 1165 1170

Ile Met Arg Lys Lys Val Ser Leu Asn Thr Tyr Thr Pro Ala Lys  
1175 1180 1185

Ile Leu Pro Thr Pro Pro Ala Thr Leu Lys Ser Thr Lys Ala Ser  
1190 1195 1200

Ser Lys Pro Ala Thr Ser Gly Pro Ser Ser Ala Val Val Pro Asn  
1205 1210 1215

Thr Ser Ser Arg Lys Lys Asn Ile Thr Ser Lys Thr Glu Thr His  
1220 1225 1230

Glu His Lys Lys Asn Thr Gly Lys Lys Lys  
1235 1240

<210> 52  
<211> 1406  
<212> PRT  
<213> Homo sapiens

<400> 52

Met Trp Ser Leu Thr Ala Ser Glu Gly Glu Ser Thr Thr Ala His Phe  
1 5 10 15

Phe Leu Gly Ala Gly Asp Glu Gly Leu Gly Thr Arg Gly Ile Gly Met  
20 25 30

Arg Pro Glu Glu Ser Asp Ser Glu Leu Leu Glu Asp Glu Glu Asp Glu  
35 40 45

Val Pro Pro Glu Pro Gln Ile Ile Val Gly Ile Cys Ala Met Thr Lys  
50 55 60

Lys Ser Lys Ser Lys Pro Met Thr Gln Ile Leu Glu Arg Leu Cys Arg  
65 70 75 80

Phe Asp Tyr Leu Thr Val Val Ile Leu Gly Glu Asp Val Ile Leu Asn  
85 90 95

Glu Pro Val Glu Asn Trp Pro Ser Cys His Cys Leu Ile Ser Phe His  
100 105 110

Ser Lys Gly Phe Pro Leu Asp Lys Ala Val Ala Tyr Ser Lys Leu Arg  
115 120 125

Asn Pro Phe Leu Ile Asn Asp Leu Ala Met Gln Tyr Tyr Ile Gln Asp  
130 135 140

Arg Arg Glu Val Tyr Arg Ile Leu Gln Glu Glu Gly Ile Asp Leu Pro  
145 150 155 160

Arg Tyr Ala Val Leu Asn Arg Asp Pro Ala Arg Pro Glu Glu Cys Asn  
165 170 175

Leu Ile Glu Gly Glu Asp Gln Val Glu Val Asn Gly Ala Val Phe Pro  
180 185 190

Lys Pro Phe Val Glu Lys Pro Val Ser Ala Glu Asp His Asn Val Tyr  
195 200 205

Ile Tyr Tyr Pro Ser Ser Ala Gly Gly Ser Gln Arg Leu Phe Arg  
210 215 220

Lys Ile Gly Ser Arg Ser Ser Val Tyr Ser Pro Glu Ser Ser Val Arg  
225 230 235 240

Lys Thr Gly Ser Tyr Ile Tyr Glu Glu Phe Met Pro Thr Asp Gly Thr  
245 250 255

Asp Val Lys Val Tyr Thr Val Gly Pro Asp Tyr Ala His Ala Glu Ala  
260 265 270

Arg Lys Ser Pro Ala Leu Asp Gly Lys Val Glu Arg Asp Ser Glu Gly  
275 280 285

Lys Glu Ile Arg Tyr Pro Val Met Leu Thr Ala Met Glu Lys Leu Val  
290 295 300

Ala Arg Lys Val Cys Val Ala Phe Lys Gln Thr Val Cys Gly Phe Asp  
305 310 315 320

Leu Leu Arg Ala Asn Gly His Ser Phe Val Cys Asp Val Asn Gly Phe  
325 330 335

Ser Phe Val Lys Asn Ser Met Lys Tyr Tyr Asp Asp Cys Ala Lys Ile  
340 345 350

Leu Gly Asn Thr Ile Met Arg Glu Leu Ala Pro Gln Phe Gln Ile Pro  
355 360 365

Trp Ser Ile Pro Thr Glu Ala Glu Asp Ile Pro Ile Val Pro Thr Thr  
370 375 380

Ser Gly Thr Met Met Glu Leu Arg Cys Val Ile Ala Ile Ile Arg His  
385 390 395 400

Gly Asp Arg Thr Pro Lys Gln Lys Met Lys Met Glu Val Lys His Pro  
405 410 415

Arg Phe Phe Ala Leu Phe Glu Lys His Gly Gly Tyr Lys Thr Gly Lys  
420 425 430

Leu Lys Leu Lys Arg Pro Glu Gln Leu Gln Glu Val Leu Asp Ile Thr  
435 440 445

Arg Leu Leu Leu Ala Glu Leu Glu Lys Glu Pro Gly Gly Glu Ile Glu  
450 455 460

Glu Lys Thr Gly Lys Leu Glu Gln Leu Lys Ser Val Leu Glu Met Tyr  
465 470 475 480

Gly His Phe Ser Gly Ile Asn Arg Lys Val Gln Leu Thr Tyr Tyr Pro  
485 490 495

His Gly Val Lys Ala Ser Asn Glu Gly Gln Asp Pro Gln Arg Glu Thr  
500 505 510

Leu Ala Pro Ser Leu Leu Leu Val Leu Lys Trp Gly Gly Glu Leu Thr  
515 520 525

Pro Ala Gly Arg Val Gln Ala Glu Glu Leu Gly Arg Ala Phe Arg Cys  
530 535 540

Met Tyr Pro Gly Gly Gln Gly Asp Tyr Ala Gly Phe Pro Gly Cys Gly  
545 550 555 560

Leu Leu Arg Leu His Ser Thr Phe Arg His Asp Leu Lys Ile Tyr Ala  
565 570 575

Ser Asp Glu Gly Arg Val Gln Met Thr Ala Ala Ala Phe Ala Lys Gly  
580 585 590

Leu Leu Ala Leu Glu Gly Glu Leu Thr Pro Ile Leu Val Gln Met Val  
595 600 605

Lys Ser Ala Asn Met Asn Gly Leu Leu Asp Ser Asp Gly Asp Ser Leu  
610 615 620

Ser Ser Cys Gln His Arg Val Lys Ala Arg Leu His His Ile Leu Gln  
625 630 635 640

Gln Asp Ala Pro Phe Gly Pro Glu Asp Tyr Asp Gln Leu Ala Pro Thr  
645 650 655

Arg Ser Thr Ser Leu Leu Asn Ser Met Thr Ile Ile Gln Asn Pro Val  
660 665 670

Lys Val Cys Asp Gln Val Phe Ala Leu Ile Glu Asn Leu Thr His Gln  
675 680 685

Ile Arg Glu Arg Met Gln Asp Pro Arg Ser Val Asp Leu Gln Leu Tyr  
690 695 700

His Ser Glu Thr Leu Glu Leu Met Leu Gln Arg Trp Ser Lys Leu Glu  
705 710 715 720

Arg Asp Phe Arg Gln Lys Ser Gly Arg Tyr Asp Ile Ser Lys Ile Pro  
725 730 735

Asp Ile Tyr Asp Cys Val Lys Tyr Asp Val Gln His Asn Gly Ser Leu  
740 745 750

Gly Leu Gln Gly Thr Ala Glu Leu Leu Arg Leu Ser Lys Ala Leu Ala  
755 760 765

Asp Val Val Ile Pro Gln Glu Tyr Gly Ile Ser Arg Glu Glu Lys Leu  
770 775 780

Glu Ile Ala Val Gly Phe Cys Leu Pro Leu Leu Arg Lys Ile Leu Leu  
785 790 795 800

Asp Leu Gln Arg Thr His Glu Asp Glu Ser Val Asn Lys Leu His Pro  
805 810 815

Leu Tyr Ser Arg Gly Val Leu Ser Pro Gly Arg His Val Arg Thr Arg  
820 825 830

Leu Tyr Phe Thr Ser Glu Ser His Val His Ser Leu Leu Ser Val Phe  
835 840 845

Arg Tyr Gly Gly Leu Leu Asp Glu Thr Gln Asp Ala Gln Trp Gln Arg  
850 855 860

Ala Leu Asp Tyr Leu Ser Ala Ile Ser Glu Leu Asn Tyr Met Thr Gln  
865 870 875 880

Ile Val Ile Met Leu Tyr Glu Asp Asn Thr Gln Asp Pro Leu Ser Glu  
885 890 895

Glu Arg Phe His Val Glu Leu His Phe Ser Pro Gly Val Lys Gly Val  
900 905 910

Glu Glu Glu Gly Ser Ala Pro Ala Gly Cys Gly Phe Arg Pro Ala Ser  
915 920 925

Ser Glu Asn Glu Glu Met Lys Thr Asn Gln Gly Ser Met Glu Asn Leu  
930 935 940

Cys Pro Gly Lys Ala Ser Asp Glu Pro Asp Arg Ala Leu Gln Thr Ser  
945 950 955 960

Pro Gln Pro Pro Glu Gly Pro Gly Leu Pro Arg Arg Ser Pro Leu Ile  
965 970 975

Arg Asn Arg Lys Ala Gly Ser Met Glu Val Leu Ser Glu Thr Ser Ser  
980 985 990

Ser Arg Pro Gly Gly Tyr Arg Leu Phe Ser Ser Ser Arg Pro Pro Thr  
995 1000 1005

Glu Met Lys Gln Ser Gly Leu Gly Phe Glu Gly Cys Ser Met Val  
1010 1015 1020

Pro Thr Ile Tyr Pro Leu Glu Thr Leu His Asn Ala Leu Ser Leu  
1025 1030 1035

Arg Gln Val Ser Glu Phe Leu Ser Arg Val Cys Gln Arg His Thr  
1040 1045 1050

Asp Ala Gln Ala Gln Ala Ser Ala Ala Leu Phe Asp Ser Met His  
1055 1060 1065

Ser Ser Gln Ala Ser Asp Asn Pro Phe Ser Pro Pro Arg Thr Leu  
1070 1075 1080

His Ser Pro Pro Leu Gln Leu Gln Gln Arg Ser Glu Lys Pro Pro  
1085 1090 1095

Trp Leu Glu Thr Arg Phe Cys His Val Gly Gln Ala Gly Leu Glu  
1100 1105 1110

Leu Leu Thr Ser Ser Asp Leu Pro Ala Ser Ala Ser Gln Ser Ala  
1115 1120 1125

Gly Ile Thr Gly Val Ser His Arg Thr Gln Pro Asp Ser Ser Gly  
1130 1135 1140

Pro Ser Ser Thr Val Ser Ser Ala Gly Pro Ser Ser Pro Thr Thr  
1145 1150 1155

Val Asp Gly Asn Ser Gln Phe Gly Phe Ser Asp Gln Pro Ser Leu  
1160 1165 1170

Asn Ser His Val Ala Glu Glu His Gln Gly Leu Gly Leu Leu Gln  
1175 1180 1185

Glu Thr Pro Gly Ser Gly Ala Gln Glu Leu Ser Ile Glu Gly Glu  
1190 1195 1200

Gln Glu Leu Phe Glu Pro Asn Gln Ser Pro Gln Val Pro Pro Met  
1205 1210 1215

Glu Thr Ser Gln Pro Tyr Glu Glu Val Ser Gln Pro Cys Gln Glu  
1220 1225 1230

Val Pro Asp Ile Ser Gln Pro Cys Gln Asp Ile Ser Glu Ala Leu  
1235 1240 1245

Ser Gln Pro Cys Gln Lys Val Pro Asp Ile Ser Gln Gln Cys Gln  
1250 1255 1260

Glu Asn His Asp Asn Gly Asn His Thr Cys Gln Glu Val Pro His  
1265 1270 1275

Ile Ser Gln Pro Cys Gln Lys Ser Ser Gln Leu Cys Gln Lys Val  
1280 1285 1290

Ser Glu Glu Val Cys Gln Leu Cys Leu Glu Asn Ser Glu Glu Val  
1295 1300 1305

Ser Gln Pro Cys Gln Gly Val Ser Val Glu Val Gly Lys Leu Val  
1310 1315 1320

His Lys Phe His Val Gly Val Gly Ser Leu Val Gln Glu Thr Leu  
1325 1330 1335

Val Glu Val Gly Ser Pro Ala Glu Glu Ile Pro Glu Glu Val Ile  
1340 1345 1350

Gln Pro Tyr Gln Glu Phe Ser Val Glu Val Gly Arg Leu Ala Gln  
1355 1360 1365

Glu Thr Ser Ala Ile Asn Leu Leu Ser Gln Gly Ile Pro Glu Ile  
1370 1375 1380

Asp Lys Pro Ser Gln Glu Phe Pro Glu Glu Ile Asp Leu Gln Ala  
1385 1390 1395

Gln Glu Val Pro Glu Glu Ile Asn  
1400 1405

<210> 53

<211> 6218

<212> DNA

<213> Homo sapiens

<400> 53

agttcgtcg agaacggagg acaccggcgg tccagggtcc tgggcagtgc cgccagagct	60
gagcggaggg cgcggcgca gaacgaatct ttgtgacatt ctctctcagc attctttatc	120
ccctgtttgc tgaagacttc gaccaaagct ggtcttagct gttggcattc tcctgagaaa	180
aggatagctt cagaaatcag aaaaacattt gggaggtgtc tagcccagtg gaccttctga	240
agagcaatgc taagaagacg tttggtttaa agaattaaaa ggaagaacaa cttaagagct	300
tcttcaaagt ttcccgcatg aaaattactt aaacttgcac acaacgtttc acaaaatctt	360
tttgtgaaaga agaaaaggaa attcagtgtg tgagtctcag caggagttaa gctaattgcag	420
cttaaaataa tgccaaaaaa gaagcgctta tctgcgggca gagtgccct gattctcttc	480
ctgtgccaga tgattagtgc actggaagta cctcttgatc caaaaacttct tgaagacttg	540
gtacagcctc caaccatcac ccaacagtct caaaaagatt acattattga ccctcgggag	600
aatattgtaa tccagtgta agccaaaggg aaaccgcccc caagctttc ctggaccggt	660
aatgggactc atttgacat cgataaagac cctctggtca ccatgaagcc tggcacagga	720
acgctcataa ttaacatcat gagcgaaggg aaagctgaga cctatgaagg agtctatcag	780
tgtacagcaa ggaacgaacg cggagctgca gtttctaata acattgttgc cggccatcc	840
agatcaccat tgtggaccaa agaaaaactt gaaccaatca cacttcaaag tggtcagtct	900
ttagtacttc cctgcagacc cccaatttggaa ttaccaccac ctataatatt ttggatggat	960

aattcctttc aaagacttcc acaaagttag agagttctc aaggttgaa tggggacctt	1020
tatTTTCCA atgtcctccc agaggacacc cgCGAAGACT atATCTGTTA TGCTAGATT	1080
aatcatactc aaaccataca gcagaagcaa CCTATTCTG TGAAGGTGAT TTCAGCTAAA	1140
tcaagttagag agaggccacc aacattttta actccagaag gcaatgcaag taacaaagag	1200
gaattaagag gaaatgtgct ttCACTGGAG TGCATTGCG AGGACTGCC TACCCCAATT	1260
atttactggg caaAGGAAGA TGGATGCTA CCCAAAACA GGACAGTTA TAAGAACATT	1320
gagaaaaacct tgCAGATCAT TCACTTTCA GAAGCAGACT CTGGAAATTa CCAATGTATA	1380
gcaaaaaatg cattaggagc catccaccat accatttctg ttAGAGTTAA AGCGGCTCCA	1440
tactggatca cAGCCCTCA AAATCTTGTG CTGTCCCCAG GAGAGGATGG GACCTTGATC	1500
TGCGAGCTA ATGGCAACCC CAAACCCAGA ATTAGCTGGT TAACAAATGG AGTCCCATA	1560
gaaattgccc CTGATGACCC CAGCAGAAAA ATAGATGGCG ATACCATTAT TTTTCAAAT	1620
gttcaagaaa gatcaagtgc AGTCTATCAG TGCAATGCCT CTAATGAATA TGGATATT	1680
CTGGCAAACG CATTGTAAGA TGTGCTGGCT GAGCACCAC GAATCCTCAC ACCTGCAAAC	1740
ACACTCTACC AGTCATTGC AACACGGCCT GCTTACTAG ACTGTGCCTT CTTGGGTCT	1800
CCTCTCCAA CCATCGAGTG GTTAAAGGA GCTAAAGGA GTGCTCTCA TGAAGATATT	1860
TATGTTTAC ATGAAAATGG AACTTTGAA ATTCTGTGG CCCAAAAGGA CAGTACAGGA	1920
ACTTATACGT GTGTTGCAAG GAATAAATTa GGGATGGCAA AGAATGAAGT TCACCTAGAA	1980
ATCAAAGATG CTACATGGAT CGTTAACAG CCCGAATATG CAGTTGTGCA AAGAGGGAGC	2040
ATGGTGTCTT TTGAATGCAA AGTGAACAT GATCACACCT TATCCCTCAC TGTCTGTGG	2100
CTGAAGGACA ACAGGAACT GCCCAGTGT GAAAGGTTCA CTGTTGACAA GGATCATCTA	2160
GTTGAGCTG ATGTCAGTGA CGATGACAGC GGGACCTACA CGTGTGTGGC CAACACCACt	2220
CTGGACAGCG TCTCCGCCAG CGCTGTGCTT AGCCTTGTG CTCCTACTCC AACTCCAGCT	2280
CCCGTTACG ATGTCCAAA TCTCCCTT GACTTAGAAC TGACAGATCA ACTTGACAAA	2340
AGTGTTCAGC TGTCAATGGAC CCCAGGCGAT GACAACAATA GCCCCATTAC AAAATTCTC	2400
ATCGAATATG AAGATGCAAT GCACAAGCCA GGGCTGTGGC ACCACCAAAC TGAAGTTCT	2460
GGAAACACAGA CCACAGCCCA GCTGAACCTG TCTCCTTACG TGAACTACTC CTTCCGCGTG	2520
ATGGCAGTGA ACAGCATTGG GAAGAGCTTG CCCAGCGAGG CGTCTGAGCA GTATTTGACG	2580
AAAGCCTCAG AACCAGATAA AAACCCACA GCTGTGGAAG GACTGGGATC AGAGCCTGAT	2640
AATTTGGTGA TTACGTGGAA GCCCTTGAAT GTTTCGAAT CTAATGGGCC AGGCCTTCAG	2700
TACAAAGTTA GCTGGCGCCA GAAAGATGGT GATGATGAAT GGACATCTGT GTTGTGGCA	2760
AATGTATCCA AATATATTGT CTCAGGCACG CCAACCTTG TTCCATAACCT GATCAAAGTT	2820

caggccctga atgacatggg gtttgcaccc gagccagctg tagtcatggg acattctgga	2880
gaagacctcc caatggtggc tcctggaaac gtgcgtgtga atgtggtgaa cagtacctta	2940
gccgaggtgc actgggaccc agtacctctg aaaagcatcc gaggacacct acaaggctat	3000
cggatttact attggaagac ccagagttca tctaaaagaa acagacgtca cattgagaaa	3060
aagatcctca cttccaagg cagcaagact catggcatgt tgccgggct agagccctt	3120
agccactaca cactgaatgt ccgagtggc aatggaaag gggagggccc agccagccct	3180
gacagagtct ttaatactcc agaaggagtc cccagtgctc cctcgcttt gaagattgtg	3240
aatccaacac tggactctct cactttggaa tggatccac cgagccaccc gaatggcatt	3300
ttgacagagt acacctaataa gtatcagcca attaacagca cacatgaatt aggcctctg	3360
gttagatttga aaattcctgc caacaagaca cggggactt taaaaaattt aaatttcagc	3420
actcgatata agtttattt ctatgcacaa acatcagcag gatcaggaag tcaaattaca	3480
gaggaagcag taacaactgt gatgaagcg atggcaagcc ggcaggtgga tattgcaact	3540
cagggctggt tcattggtct gatgtgtgct gttgctctcc ttatcttaat tttgctgatt	3600
gtttgcttca tcagaagaaa caagggtggt aaatatccag taaaagaaaa ggaagatgcc	3660
catgctgacc ctgaaatcca gcctatgaag gaagatgatg ggacatttg agaatacagt	3720
gatgcagaag accacaagcc tttgaaaaaa ggaagtcgaa ctccttcaga caggactgt	3780
aaaaaagaag atagtgacga cagcctagtt gactatggag aaggggttaa tggccagttc	3840
aatgaggatg gtcctttat tggacaatac agtggtaaga aagagaaaga gccggctgaa	3900
ggaaacgaaa gtcagaggc accttctcct gtcaacgcca tgaattcctt tgttaattt	3960
ttaagcttctt tgccaatatt ccatttctct agaatgttta tcctaagcac ttgttgtca	4020
gccctctcat actatgaaca tatgggtaga gagtatattt tctgctgtat gttgttattt	4080
tgagaatagt tacagcaaaa acataactca gtcaaattgt atgttaatataa gaactggaaat	4140
gcaaagtgca tacttttca ttcaaaatgg gtattcttga tttcctcaga actgataaaaa	4200
aataatgcaa catcaccaac agatcctgtt atttcctctg caggatacag ttcaatatga	4260
tgcataaaaa atgctccaca tttaaaggac ataccgtgt atgttatgaa aacatggttt	4320
gatactttgt ttatactacc ctcagctgaa cccctatata tgaattccgt tttcattgtc	4380
aagagtgtta ctgttagtatt ctctagaact tcaatgtctt tgtggacatt gttgtgaaat	4440
tggtgactat gtatactgtgt cgtagtctt tttggagac tgtaggaac agtttgcata	4500
gtatatactt gctaaatgag ttcattatga cagtcacatt gctgatgctt actgagaact	4560
attacctact cttggctcct gttactccgt aggcttctta atcttccagg cattacagca	4620
gcacagtgtt ctactttta catcatttct atgttcgggtt gtttttaggc ataaacaatg	4680

tgtattgcag	tgcatttcgg	catttgcgc	atactgaaag	aatcaaaaac	aatcatcca	4740
aattaaattt	caaacattat	ttcagagaac	acagggcaag	acacatacag	tgccttcaga	4800
tattaaggcat	tccacaacat	cgtgcattct	gtatcagctg	gtccagtcca	ttctgggtcc	4860
tagattactg	tcattgtcta	aaagtaactt	ttaaaaagca	gagttcatga	aaactgcaat	4920
gctggggaaaa	gaaggaaaaca	tgaaaataaa	aataagacag	tttattagaa	atagcatttc	4980
ctcataagca	taaaaagaaa	atctttgttg	ccaaactgaag	cacatgatga	ttttgtggtc	5040
ctttatgggtt	tctattacat	gcagtaagaa	agatgtcaac	atgctagaaa	attaatttttta	5100
aaactaagtt	attccaacac	taaaagcata	caacagcatg	ccaaacagtaa	tatattatttc	5160
tccaaagactt	tacctatgta	agtgttcaaa	actctgcagc	attaaacaac	gtgtatgcaa	5220
attgttatgg	atacatttca	gaatctaaga	aatcaggcaa	gtgcttaaaa	ggccaacgggt	5280
ccaaagggatt	acatctgcag	tttaaaaagt	aaatatataat	tctatcgat	tcataaacaa	5340
tatctatcaa	atgggttacc	tccaaatatg	aaaatctata	acaacctatg	gttgaaggaa	5400
tgctcagttt	catttgccaa	taaattgggtt	tctcataact	tgcatcaagt	ttaattttaa	5460
gtaaagcttt	ttatatgttag	atattttgtt	gaatttgtaa	atacacttaa	aatgttagatg	5520
ctatatgctt	ataggtgtta	catacaaata	aacatgcaat	gtttatgttg	tactgtataa	5580
gaggtaagct	aattaatgca	gtgaatggga	ttggaaagca	tctacttaaa	tatctattgg	5640
gttccccccct	cccccccacct	ttttgctgt	gaaactgaaa	tagtgaactt	ttctacgtat	5700
tgacagcaga	tttttcgatg	aaatcttcag	agctttgcct	atggggcaca	gtaggcctag	5760
taacctggca	tgtttgatat	atgttagttaa	agcataattt	aaagtaatcc	caggtaaaga	5820
tggccctaaa	tactttcatg	tctctatatt	cattttcac	agatccacct	gtctcttgaa	5880
aatataaaaa	gacaaaacag	gtttgccttg	gcatcagaga	gcacaaaagat	taaaagttac	5940
tttaaatttg	ccaatatttt	gggagaacaa	taaaactaca	tttttcctc	ttccatactg	6000
gtagatgcga	aatttatctg	tgcatgaaag	ggtcacttct	gtaatagtgc	aacagatttgc	6060
gtattaaaaaa	ttaaatgtgg	ttttaaaagt	tcctctctct	tttgtaattt	atgttcccaa	6120
ttgagtgtga	atgtccaagt	aatggtgtat	gtaatggtac	aggcaaatgt	gactggattt	6180
ccctcaaaaaa	agtaacttattt	aaacagtctt	gatctttt			6218

<210> 54  
 <211> 1180  
 <212> PRT  
 <213> Homo sapiens

<400> 54

Met Gln Leu Lys Ile Met Pro Lys Lys Lys Arg Leu Ser Ala Gly Arg  
 1 . 5 . 10 . 15

Val Pro Leu Ile Leu Phe Leu Cys Gln Met Ile Ser Ala Leu Glu Val  
20 25 30

Pro Leu Asp Pro Lys Leu Leu Glu Asp Leu Val Gln Pro Pro Thr Ile  
35 40 45

Thr Gln Gln Ser Pro Lys Asp Tyr Ile Ile Asp Pro Arg Glu Asn Ile  
50 55 60

Val Ile Gln Cys Glu Ala Lys Gly Lys Pro Pro Pro Ser Phe Ser Trp  
65 70 75 80

Thr Arg Asn Gly Thr His Phe Asp Ile Asp Lys Asp Pro Leu Val Thr  
85 90 95

Met Lys Pro Gly Thr Gly Thr Leu Ile Ile Asn Ile Met Ser Glu Gly  
100 105 110

Lys Ala Glu Thr Tyr Glu Gly Val Tyr Gln Cys Thr Ala Arg Asn Glu  
115 120 125

Arg Gly Ala Ala Val Ser Asn Asn Ile Val Val Arg Pro Ser Arg Ser  
130 135 140

Pro Leu Trp Thr Lys Glu Lys Leu Glu Pro Ile Thr Leu Gln Ser Gly  
145 150 155 160

Gln Ser Leu Val Leu Pro Cys Arg Pro Pro Ile Gly Leu Pro Pro Pro  
165 170 175

Ile Ile Phe Trp Met Asp Asn Ser Phe Gln Arg Leu Pro Gln Ser Glu  
180 185 190

Arg Val Ser Gln Gly Leu Asn Gly Asp Leu Tyr Phe Ser Asn Val Leu  
195 200 205

Pro Glu Asp Thr Arg Glu Asp Tyr Ile Cys Tyr Ala Arg Phe Asn His  
210 215 220

Thr Gln Thr Ile Gln Gln Lys Gln Pro Ile Ser Val Lys Val Ile Ser  
225 230 235 240

Ala Lys Ser Ser Arg Glu Arg Pro Pro Thr Phe Leu Thr Pro Glu Gly  
245 250 255

Asn Ala Ser Asn Lys Glu Glu Leu Arg Gly Asn Val Leu Ser Leu Glu  
260 265 270

Cys Ile Ala Glu Gly Leu Pro Thr Pro Ile Ile Tyr Trp Ala Lys Glu  
275 280 285

Asp Gly Met Leu Pro Lys Asn Arg Thr Val Tyr Lys Asn Phe Glu Lys  
290 295 300

Thr Leu Gln Ile Ile His Val Ser Glu Ala Asp Ser Gly Asn Tyr Gln  
305 310 315 320

Cys Ile Ala Lys Asn Ala Leu Gly Ala Ile His His Thr Ile Ser Val  
325 330 335

Arg Val Lys Ala Ala Pro Tyr Trp Ile Thr Ala Pro Gln Asn Leu Val  
340 345 350

Leu Ser Pro Gly Glu Asp Gly Thr Leu Ile Cys Arg Ala Asn Gly Asn  
355 360 365

Pro Lys Pro Arg Ile Ser Trp Leu Thr Asn Gly Val Pro Ile Glu Ile  
370 375 380

Ala Pro Asp Asp Pro Ser Arg Lys Ile Asp Gly Asp Thr Ile Ile Phe  
385 390 395 400

Ser Asn Val Gln Glu Arg Ser Ser Ala Val Tyr Gln Cys Asn Ala Ser  
405 410 415

Asn Glu Tyr Gly Tyr Leu Leu Ala Asn Ala Phe Val Asn Val Leu Ala  
420 425 430

Glu Pro Pro Arg Ile Leu Thr Pro Ala Asn Thr Leu Tyr Gln Val Ile  
435 440 445

Ala Asn Arg Pro Ala Leu Leu Asp Cys Ala Phe Phe Gly Ser Pro Leu  
450 455 460

Pro Thr Ile Glu Trp Phe Lys Gly Ala Lys Gly Ser Ala Leu His Glu  
465 470 475 480

Asp Ile Tyr Val Leu His Glu Asn Gly Thr Leu Glu Ile Pro Val Ala  
485 490 495

Gln Lys Asp Ser Thr Gly Thr Tyr Thr Cys Val Ala Arg Asn Lys Leu  
500 505 510

Gly Met Ala Lys Asn Glu Val His Leu Glu Ile Lys Asp Ala Thr Trp  
515 520 525

Ile Val Lys Gln Pro Glu Tyr Ala Val Val Gln Arg Gly Ser Met Val  
530 535 540

Ser Phe Glu Cys Lys Val Lys His Asp His Thr Leu Ser Leu Thr Val  
545 550 555 560

Leu Trp Leu Lys Asp Asn Arg Glu Leu Pro Ser Asp Glu Arg Phe Thr  
565 570 575

Val Asp Lys Asp His Leu Val Val Ala Asp Val Ser Asp Asp Asp Ser  
580 585 590

Gly Thr Tyr Thr Cys Val Ala Asn Thr Thr Leu Asp Ser Val Ser Ala  
595 600 605

Ser Ala Val Leu Ser Val Val Ala Pro Thr Pro Thr Pro Ala Pro Val  
610 615 620

Tyr Asp Val Pro Asn Pro Pro Phe Asp Leu Glu Leu Thr Asp Gln Leu  
625 630 635 640

Asp Lys Ser Val Gln Leu Ser Trp Thr Pro Gly Asp Asp Asn Asn Ser  
645 650 655

Pro Ile Thr Lys Phe Ile Ile Glu Tyr Glu Asp Ala Met His Lys Pro  
660 665 670

Gly Leu Trp His His Gln Thr Glu Val Ser Gly Thr Gln Thr Thr Ala  
675 680 685

Gln Leu Asn Leu Ser Pro Tyr Val Asn Tyr Ser Phe Arg Val Met Ala  
690 695 700

Val Asn Ser Ile Gly Lys Ser Leu Pro Ser Glu Ala Ser Glu Gln Tyr  
705 710 715 720

Leu Thr Lys Ala Ser Glu Pro Asp Lys Asn Pro Thr Ala Val Glu Gly  
725 730 735

Leu Gly Ser Glu Pro Asp Asn Leu Val Ile Thr Trp Lys Pro Leu Asn  
740 745 750

Gly Phe Glu Ser Asn Gly Pro Gly Leu Gln Tyr Lys Val Ser Trp Arg  
755 760 765

Gln Lys Asp Gly Asp Asp Glu Trp Thr Ser Val Val Val Ala Asn Val  
770 775 780

Ser Lys Tyr Ile Val Ser Gly Thr Pro Thr Phe Val Pro Tyr Leu Ile  
785 790 795 800

Lys Val Gln Ala Leu Asn Asp Met Gly Phe Ala Pro Glu Pro Ala Val  
805 810 815

Val Met Gly His Ser Gly Glu Asp Leu Pro Met Val Ala Pro Gly Asn  
820 825 830

Val Arg Val Asn Val Val Asn Ser Thr Leu Ala Glu Val His Trp Asp  
835 840 845

Pro Val Pro Leu Lys Ser Ile Arg Gly His Leu Gln Gly Tyr Arg Ile  
850 855 860

Tyr Tyr Trp Lys Thr Gln Ser Ser Ser Lys Arg Asn Arg Arg His Ile  
865 870 875 880

Glu Lys Lys Ile Leu Thr Phe Gln Gly Ser Lys Thr His Gly Met Leu  
885 890 895

Pro Gly Leu Glu Pro Phe Ser His Tyr Thr Leu Asn Val Arg Val Val  
900 905 910

Asn Gly Lys Gly Glu Gly Pro Ala Ser Pro Asp Arg Val Phe Asn Thr  
915 920 925

Pro Glu Gly Val Pro Ser Ala Pro Ser Ser Leu Lys Ile Val Asn Pro  
930 935 940

Thr Leu Asp Ser Leu Thr Leu Glu Trp Asp Pro Pro Ser His Pro Asn  
945 950 955 960

Gly Ile Leu Thr Glu Tyr Thr Leu Lys Tyr Gln Pro Ile Asn Ser Thr  
965 970 975

His Glu Leu Gly Pro Leu Val Asp Leu Lys Ile Pro Ala Asn Lys Thr  
980 985 990

Arg Trp Thr Leu Lys Asn Leu Asn Phe Ser Thr Arg Tyr Lys Phe Tyr  
995 1000 1005

Phe Tyr Ala Gln Thr Ser Ala Gly Ser Gly Ser Gln Ile Thr Glu  
1010 1015 1020

Glu Ala Val Thr Thr Val Asp Glu Ala Met Ala Ser Arg Gln Val  
1025 1030 1035

Asp Ile Ala Thr Gln Gly Trp Phe Ile Gly Leu Met Cys Ala Val  
1040 1045 1050

Ala Leu Leu Ile Leu Ile Leu Leu Ile Val Cys Phe Ile Arg Arg  
1055 1060 1065

Asn Lys Gly Gly Lys Tyr Pro Val Lys Glu Lys Glu Asp Ala His  
1070 1075 1080

Ala Asp Pro Glu Ile Gln Pro Met Lys Glu Asp Asp Gly Thr Phe  
1085 1090 1095

Gly Glu Tyr Ser Asp Ala Glu Asp His Lys Pro Leu Lys Lys Gly  
1100 1105 1110

Ser Arg Thr Pro Ser Asp Arg Thr Val Lys Lys Glu Asp Ser Asp  
1115 1120 1125

Asp Ser Leu Val Asp Tyr Gly Glu Gly Val Asn Gly Gln Phe Asn  
1130 1135 1140

Glu Asp Gly Ser Phe Ile Gly Gln Tyr Ser Gly Lys Lys Glu Lys  
1145 1150 1155

Glu Pro Ala Glu Gly Asn Glu Ser Ser Glu Ala Pro Ser Pro Val  
1160 1165 1170

Asn Ala Met Asn Ser Phe Val  
1175 1180